

COMPOSITIONS AND METHODS FOR INCREASING AMINO ACID ABSORPTION IN MAMMALS

Cross-Reference to Related Applications

5 This application claims the benefit of U.S. Provisional Application Serial
Number 60/273,263, filed March 2, 2001, under 35 U.S.C. 119(e) and U.S.
Provisional Application Serial Number 60/344,088, filed December 26, 2001, under
35 U.S.C. 119(e).

Background of the Invention

10 In dogs, it is thought that the ability to absorb essential amino acids such as
tryptophan and leucine may be limiting to cellular metabolism. Recent research
designed to characterize the amino acid absorption capacity of the brush border
(lumen facing) membranes of dog enterocytes suggests that peptide absorption may
15 be particularly important given the relatively low amount of free amino acid
transport capacity that was observed. Buddington RK, Paulsen DB. Development
of the Canine and Feline Gastrointestinal Tract. In: Reinhart GA, Carey DP, eds.
*Recent Advances in Canine and Feline Nutrition, Vol. II: 1998 Iams Nutrition
Symposium Proceedings*. Wilmington: Orange Frazer Press, 1998; 195-215. Data
20 collected from studies designed to understand the quantitative importance of free
versus peptide amino acids in other monogastric animals strongly indicates that
peptide-bound amino acids account for the majority of amino acids absorbed by
enterocytes from the intestinal lumen (Matthews, DM. *Protein Absorption,
Development and Present State of the Subject*, New York: Wiley-Liss, 1991.) and
25 that the rate of peptide-derived amino acid absorption is faster than that by
equivalent amounts of free amino acids. Ohkohchi N, Andoh T, Ohi R, Mori S.
Defined formula diets alter characteristics of the intestinal transport of amino acid
and peptide in growing rats. *J Pediatr Gastroenterol Nutr* 1990 May; 10(4):490-6.

30 Two types of peptide transporters have been cloned from monogastric
animals. Liang R, Fei YJ, Prasad PD, Ramamoorthy S, Han H, Yang-Feng TL,

Hediger MA, Ganapathy V, Leibach FH. Human intestinal H⁺/peptide cotransporter. Cloning, functional expression, and chromosomal localization. *J Biol Chem* 1995 Mar 24; 270(12):6456-63. Liu W, Liang R, Ramamoorthy S, Fei YJ, Ganapathy ME, Hediger MA, Ganapathy V, Leibach FH. Molecular cloning of PEPT 2, a new member of the H⁺/peptide cotransporter family, from human kidney. *Biochim Biophys Acta* 1995 May 4; 1235(2):461-6. PepT1 is an H⁺-dependent, low-affinity (mM), high-velocity, transporter that is predominately localized primarily to the brush border membranes of mature enterocytes of intestinal villi. PepT2 is an H⁺-dependent, high-affinity (μM), low-velocity, transporter that is expressed in the greatest abundance in the apical membranes of renal proximal tubular epithelial cells. An important feature of the peptide transporters is their ability to recognize and transport most di- and tripeptides, albeit with a range of relative affinities for different peptides. In addition, both transporters recognize the β-lactam antibiotics, and carboxyl-terminal modified free amino acids. The physiologic functions of these transporters are thought to be to absorb di- and tripeptides from the digesta and from the blood, respectively. Although molecular evidence has not been acquired, there is strong biochemical evidence for a different peptide transport protein that functions in the basolateral membrane of these cells. Saito H, Inui KI. Dipeptide transporters in apical and basolateral membranes of the human intestinal cell line Caco-2. *Am J Physiol* 1993 Aug; 265(2 Pt 1):G289-94. Thwaites DT, Brown CD, Hirst BH, Simmons NL. Transepithelial glycylsarcosine transport in intestinal Caco-2 cells mediated by the expression of H⁺-coupled carriers at both the apical and basal membranes. *J Biol Chem* 1993 Apr 15; 268(11):7640-2.

Research with Caco-2 cells indicates that PepT1 transporter mRNA, protein, and activity increases in a manner consistent with a direct effect of increased extracellular substrate concentrations. Walker D, Thwaites DT, Simmons NL, Gilbert HJ, Hirst BH. Substrate upregulation of the human small intestinal peptide transporter, hPepT1. *J Physiol* 1998 Mar 15; 507(Pt 3):697-706. In contrast to mRNAs for essential amino acid transporters, intestinal studies show that the

expression of peptide transporter mRNA increases in response to increased dietary protein. Erickson RH, Gum JR Jr, Lindstrom MM, McKean D, Kim YS. Regional expression and dietary regulation of rat small intestinal peptide and amino acid transporter mRNAs. *Biochem Biophys Res Commun* 1995 Nov 2; 216(1):249-57.

- 5 Similarly, expression in intestinal mucosa of PepT1 mRNA and protein increases in response to tissue trauma, whereas the mRNA for essential amino acid transporters decreases. Tanaka H, Miyamoto KI, Morita K, Haga H, Segawa H, Shiraga T, Fujioka A, Kuoda T, Taketani Y, Hisano S, Fukui Y, Kitagawa K, Takeda E. Regulation of the PepT1 peptide transporter in the rat small intestine in response to
10 5-fluorouracil-induced injury. *Gastroenterology* 1998 Apr; 114(4):714-23.

- Few studies have been conducted to evaluate the potential for the dog to absorb quantitatively significant amounts of essential amino acids in the form of small peptides, and whether this capacity can be regulated by substrate supply. Accordingly, there is still a need to evaluate the potential for the absorption of
15 peptide-bound leucine and tryptophan by putative canine peptide transporters. It would thus be desirable to provide the nucleic acid sequence encoding canine PepT1. It would also be desirable to provide mRNA transcripts corresponding to cPepT1. It would further be desirable to characterize the function of cPepT1 by GlySar uptake and identify di- and tripeptides well recognized by cPepT1, as well as
20 characterize the effect of supplemental peptide substrate on the transport capacity of canine PepT1 (cPepT1).

Summary of the Invention

- The present invention provides novel isolated and purified nucleic acids
25 (RNA or DNA) encoding, or complementary to, canine PepT1 (cPepT1). The nucleic acid may be SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:9, or SEQ ID NO:20 or may be a nucleic acid that hybridizes under moderate or stringent hybridization conditions to any of these sequences. Also provided are peptides encoded by these nucleic acids, such as SEQ ID NO:13 or SEQ ID NO:21.

The present invention also provides a method for determining canine PepT1-transportability of a peptide, or method for determining a peptide with beneficial nutritional property in an animal, comprising providing an immortalized kidney distal tubule epithelial (Madin-Darby Canine Kidney (MDCK)) cell and a peptide
5 having 2 to 10 amino acids, and determining the amount of the peptide transported into the cell, wherein the amount correlates with the canine PepT1-transportability of the peptide. A peptide with beneficial nutritional properties in an animal is a peptide that contains at least one essential amino acid that is absorbed at a rate higher than the rate of the amino acid if it were free rather than in a peptide-bound
10 form. The peptide may be a dipeptide, tripeptide, or tetrapeptide such as, for example, GlySar, GlyGly, AlaHis, β -AlaHis (carnosine), GlnGln, GlyMet, LeuMet, LeuTrp, MetLeu, MetMet, MetPhe, MetPro, TrpLeu, TrpTrp, GlnGlu, MetGlu, MetLys, TrpGly, MetGlyMetMet (SEQ ID NO:10), TrpGlyGly, LeuArg, ArgLeu, GlyLeu, or ArgTrp. The cell used in the method may be in medium at a pH of
15 between about 5 and 8; or at a pH of about 5.5 to 7.5, or even at about 6 to 6.5. The peptide may be present at a concentration of about 10 nM to about 50 mM.

The characterization of GlySar uptake by immortalized MDCK cells demonstrates that MDCK cells express PepT1-like activity, confirming detection of PepT1 mRNA expression by MDCK cells and the use of MDCK cells as a model to
20 characterize the biochemical function of canine PepT1.

The cPepT1 of the present invention is also capable of recognizing a variety of di- and tripeptides, including those that contain the essential amino acids leucine and tryptophan, considered to be of especial importance to canine nutrition. In addition, H⁺-dependent peptide transport in cultured MDCK cells can be stimulated
25 by at least two of PepT1 substrates, GlySar and carnosine. Moreover, H⁺-dependent uptake of GlySar by MDCK is sensitive to nutrient deprivation and Insulin-like Growth factor I (IGF-I).

The present invention further provides a dietary composition with improved nutritional benefit for an animal comprising at least one peptide identified by the
30 method described above.

The present invention provides a process for altering the absorption of essential amino acids in an animal, such as a dog, comprising the steps of feeding the animal a diet containing the dietary composition described above; and maintaining the animal on the diet for a sufficient period of time to allow the composition to be absorbed by the digestive system of the animal. The diet may comprise about 20 to about 30% crude protein, about 10 to about 20% fat, and about 3 to about 10% dietary fiber.

As used herein, the term "cPepT1" includes variants or biologically active or inactive fragments of this transport protein. A "variant" of the polypeptide is a cPepT1 protein that is not completely identical to a native cPepT1 protein. A variant cPepT1 protein can be obtained by altering the amino acid sequence by insertion, deletion or substitution of one or more amino acid. The amino acid sequence of the protein is modified, for example by substitution, to create a polypeptide having substantially the same or improved qualities as compared to the native polypeptide. The substitution may be a conserved substitution. A "conserved substitution" is a substitution of an amino acid with another amino acid having a similar side chain. A conserved substitution would be a substitution with an amino acid that makes the smallest change possible in the charge of the amino acid or size of the side chain of the amino acid (alternatively, in the size, charge or kind of chemical group within the side chain) such that the overall peptide retains its spacial conformation but has altered biological activity. For example, common conserved changes might be Asp to Glu, Asn or Gln; His to Lys, Arg or Phe; Asn to Gln, Asp or Glu and Ser to Cys, Thr or Gly. Alanine is commonly used to substitute for other amino acids. The 20 common amino acids can be grouped as follows: alanine, valine, leucine, isoleucine, proline, phenylalanine, tryptophan and methionine having nonpolar side chains; glycine, serine, threonine, cystine, tyrosine, asparagine and glutamine having uncharged polar side chains; aspartate and glutamate having acidic side chains; and lysine, arginine, and histidine having basic side chains. Stryer, L. *Biochemistry* (2d edition) W. H. Freeman and Co. San Francisco (1981), p. 14-15; Lehninger, A. *Biochemistry* (2d ed., 1975), p. 73-75. It

is known to those of skill in the art that transport of other, less common, amino acids such as hydroxylysine, or derivatives of any one of the 20 common amino acids listed above would also be within the scope of this invention.

It is known that variant polypeptides can be obtained based on substituting certain amino acids for other amino acids in the polypeptide structure in order to modify or improve biological activity. For example, through substitution of alternative amino acids, small conformational changes may be conferred upon a polypeptide that result in increased bioactivity. Alternatively, amino acid substitutions in certain polypeptides may be used to provide residues that may then be linked to other molecules to provide peptide-molecule conjugates that retain sufficient properties of the starting polypeptide to be useful for other purposes.

One can use the hydropathic index of amino acids in conferring interactive biological function on a polypeptide, wherein it is found that certain amino acids may be substituted for other amino acids having similar hydropathic indices and still retain a similar biological activity. Alternatively, substitution of like amino acids may be made on the basis of hydrophilicity. It is noted that substitutions can be made based on the hydrophilicity assigned to each amino acid. In using either the hydrophilicity index or hydropathic index, which assigns values to each amino acid, it is preferred to conduct substitutions of amino acids where these values are ± 2 , with ± 1 being particularly preferred, and those with in ± 0.5 being the most preferred substitutions.

The variant cPepT1 protein comprises at least seven amino acid residues, preferably about 20 to about 700 residues, and more preferably about 50 to about 700 residues, wherein the variant cPepT1 protein has at least 50%, preferably at least about 80%, and more preferably at least about 90% but less than 100%, contiguous amino acid sequence homology or identity to the amino acid sequence of a corresponding native cPepT1 protein.

The amino acid sequence of the variant cPepT1 protein corresponds essentially to the native cPepT1 protein amino acid sequence. As used herein “correspond essentially to” refers to a polypeptide sequence that will elicit an

absorption value substantially the same as the absorption stimulated by native cPepT1 protein. Such absorption may be at least 60% of the level generated by native cPepT1 protein, and may even be at least 80% of the level generated by native cPepT1 protein.

5 A variant of the invention may include amino acid residues not present in the corresponding native cPepT1 protein, or may include deletions relative to the corresponding native cPepT1 protein. A variant may also be a truncated “fragment” as compared to the corresponding native cPepT1 protein, *i.e.*, only a portion of a full-length protein. cPepT1 protein variants also include peptides having at least
10 one D-amino acid.

The cPepT1 protein of the present invention may be expressed from an isolated nucleic acid (DNA or RNA) sequence encoding the cPepT1 protein. Amino acid changes from the native to the variant cPepT1 protein may be achieved by changing the codons of the corresponding nucleic acid sequence. “Recombinant” is
15 defined as a peptide or nucleic acid produced by the processes of genetic engineering. It should be noted that it is well-known in the art that, due to the redundancy in the genetic code, individual nucleotides can be readily exchanged in a codon, and still result in an identical amino acid sequence. The terms “protein,” “peptide” and “polypeptide” are used interchangeably herein.

20

Brief Description of the Figures

Figure 1 is a photograph of an electrophoresis gel showing the partial length canine PepT1 cDNA reaction products generated by reverse transcription-
polymerase chain reaction (RT-PCR) methodology. Partial length canine PepT1
25 (cPepT1, about 783 bp) cDNAs were generated by reverse transcription-polymerase chain reaction (RT-PCR) methodology. RT-PCR reaction products were generated using mRNA isolated from canine jejunal epithelium and two different PCR primer sets. Gel contents are as follows: lane 1, 1 Kb molecular weight DNA ladder; lane
2, negative control PCR reaction (lacks Taq polymerase); lane 3, PCR reaction
30 products using primer set 4 (corresponding to base pairs 83 to 863 of rabbit PepT1);

lane 4, PCR reaction products using primer set 10~780 bp cDNA product using primer set 10 (corresponding to base pairs 85 to 861 of rabbit PepT1). Note the reaction products in lanes 3 and 4 of about 780 base pairs.

Figures 2A and 2B are photographs of agarose gels showing the representative results of restriction analyses of pCR®II/cPepT1 plasmids generated by TA-cloning of primer set 4-derived RT-PCR cDNA. Restriction analyses of pCR®II/cPepT1 plasmids generated by TA-cloning of primer set 4-derived RT-PCR cDNA are shown in these figures. Data are representative of four cDNA-containing plasmids from a total of fifty-six “positive” bacterial colonies selected by blue/white screening. TA-clones were amplified, pCR®II/cDNA vectors isolated, and Xho I and Kpn I endonucleases restriction products size-separated through 1.2% agarose gels. **Figure 2A** is a photograph of an agarose gel showing representative results of the PCR-based analyses of TA-clone 26. In particular, analyses of pCR®II/cPepT1-26 (TA-clone 26) are shown; lane 1, 1 DNA size standard; lane 2, minus endonuclease-restriction control (uncut pCR®II plasmid); lane 3, positive restriction control (Xho I restriction of empty pCR®II vector); lane 4, uncut pCR®II/cPepT1-26 (Clone 26); lane 5, Xho I- and Kpn I-restricted Clone 26. Note that empty pCR®II vector is 3.9 kb in size and that lane 5 contains a product of about 780 bp. **Figure 2B** is a photograph of an agarose gel showing representative results of the PCR-based analyses of TA-clone 4 and 6. In particular, analyses of TA-clone 4 and 6 are shown; lane 1, DNA size standard; lane 2, uncut pCR®II/cPepT1-4 (TA-clone 4); lane 3, Xho I- and Kpn I-restricted pCR®II/cPepT1-4; lane 4, uncut pCR®II/cPepT1-6 (TA-clone 6); lane 5, Xho I- and Kpn I-restricted pCR®II/cPepT1-6. Note that empty pCR®II vector is 3.9 kb in size and that lane 3 does not contain a product of about 780 bp, whereas lane 5 does.

Figures 3A and 3B are photographs showing the representative results of Northern blot identification of cPepT1 mRNA expression by canine tissues and MDCK cells using canine intestinal epithelium-derived RT-PCR cDNA. Arrangement of RNA isolated from tissue or cell homogenates on both blots is as follows: lane 1, kidney (animal #1031A); lane 2, kidney (animal K-9-1); lane 3,

MDCK cells; lane 4, jejunal epithelium (animal K-9-4). **Figure 3A** is a photograph showing the Northern blot identification of A⁺RNA (3 µg/lane) that was hybridized with [³²P]-cPepT1-26 cDNA. **Figure 3B** is a photograph showing the Northern blot identification of total RNA (20 µg/lane) that was hybridized with [³²P]-cPepT1-6 cDNA.

Figure 4 is a photograph showing the representative results of Northern blot identification of cPepT1 mRNA expression in canine tissues using full-length rabbit PepT1 cDNA. Ten µg total RNA (lane 1) or 6 µg A⁺RNA (lanes 2 to 5) were isolated from liver and kidney tissues from three animals. Lane 1, liver (animal #1042A); lane 2, liver (animal #1008A); lane 3, kidney (animal #1008A); lane 4, liver (animal #1031A); lane 5, kidney (animal #1031A).

Figure 5 is the partial-length nucleic acid sequence of canine PepT1 cDNA of the present invention that was cloned from MDCK cells (SEQ ID NO:9). The 381 base pairs of TA clone PepT1-6R-20 shares 79% homology to base pairs 259 to 640 of rabbit PepT1 (GenBank acc no. 473375).

Figure 6 is a graph illustrating the influence of extracellular GlySar concentrations on GlySar uptake by confluent MDCK cells in pH 6.0 media. By graphical evaluation, an apparent K_m of about 4 mM was demonstrated. Each data point is the mean of 5 to 6 observations and all coefficients of variation were less than 15%.

Figure 7 is a graph illustrating the protein content of MDCK cells cultured in DMEM or LHM. Values are the means ± SD of protein content of wells (n = 12) of MDCK cells after seeding at 60,000 or 120,000 cells/well, culture for 1 d in DMEM, and then culture in DMEM or LHM for 1, 2, 3, or 5 d (Days 2, 3, 4 and 6, respectively). Protein content was determined by the method of Lowry, using bovine serum albumin as the standard.

Figure 8 is a graph illustrating GlySar (2.88 µM) uptake in pH 6.0 or pH 7.4 buffer by MDCK cells cultured in DMEM or LHM. Uptake was measured in the absence (pH 7.4) or presence (pH 6.0) of an extracellular-to-intracellular H⁺ gradient.

Figure 9 is a graph illustrating H^+ -dependent [3H]-GlySar (2.88 μM) uptake by MDCK cells cultured in DMEM or LHM. Values were calculated as the difference in GlySar uptake in the presence (pH 6.0 uptake buffer) and absence (pH 7.4 uptake buffer) of an extracellular-to-intercellular H^+ proton gradient.

5 **Figure 10** is a graph illustrating pH-dependent GlySar uptake by MDCK cells seeded at 60,000 cells/well and cultured in LHM for 2 days. pH-dependent GlySar (2.88 μM) uptake by MDCK cells cultured with standard conditions. Values represent the H^+ -dependent GlySar uptake means \pm SD of wells ($n = 16$) of MDCK cells, calculated as the difference from GlySar uptake in the presence of pH 6.0 or
10 7.4 buffers.

Figure 11 is a graph illustrating the effect of time on GlySar uptake (100 μM) by MDCK cells. By-minute time course for GlySar (uptake by MDCK cells cultured with standard conditions. Mean \pm SD GlySar uptake wells of cells ($n = 6$) were assayed at 3.75, 7.5, 15, 30, 60, or 120 min.

15 **Figure 12** is a graph illustrating the effect of GlySar concentration on MDCK cells seeded at 60K/well grown in LHM. The graph indicates the K_m characterization (1.0 mM) of H^+ -dependent GlySar uptake by MDCK cells. Each value represents the mean \pm SD uptake of GlySar by wells ($n = 8$) of MDCK cells cultured using standard conditions.

20 **Figure 13** is a graph illustrating the inhibition of peptide uptake by MDCK cells with antibiotics. The mean \pm SD are the uptake of GlySar by wells ($n = 5-8$) of MDCK cells in the absence or presence of GlySar (1 mM) Penicillin-G (3 mM), cefadroxil (30 μM), or cefadroxil (3 mM).

Figure 14 is a graph illustrating the inhibition of peptide uptake by MDCK
25 cells with Gly-containing peptides. The mean \pm SD uptake of GlySar by wells ($n = 7-8$) of MDCK cells in the absence or presence of indicated competitor substrates (1 mM).

Figure 15 is a graph illustrating the inhibition of 100 μM GlySar uptake by 1mM TrpLeu, LeuTrp, Leu, or Trp in the absence (pH 7.5) and presence (no pH

designation) of a proton gradient and 1 mM of indicated substrates. Values are the mean \pm SD uptake of GlySar by wells (n = 7-8) of MDCK cells.

Figure 16 is a graph illustrating the inhibition of 100 μ M GlySar uptake by MDCK cells in the absence (pH 7.5) and presence (no pH designation) of a proton
5 gradient and 1 mM of Trp-containing peptides. Values are the mean \pm SD uptake of GlySar by wells (n = 7-8) of MDCK cells.

Figure 17 is a graph illustrating the inhibition of 100 μ M GlySar uptake by MDCK cells in the absence (pH 7.5) and presence (no pH designation) of a proton
10 gradient and 100 μ M of Trp-containing peptides. Values are the mean \pm SD uptake of GlySar by wells (n = 8) of MDCK cells.

Figure 18 is a graph illustrating the IC₅₀ inhibition of H⁺-dependent GlySar uptake by TrpLeu and TrpTrp. K_i values were determined for inhibition of H⁺-
15 dependent 100 μ M GlySar uptake by MDCK cells in the presence of 0, 0.025, 0.1, 0.4, or 1.6 mM TrpTrp or TrpLeu. Values are the mean \pm SD uptake of GlySar by wells (n = 6-8) of MDCK cells.

Figure 19 is a graph illustrating substrate (10mM) regulation of protein content of MDCK cells cultured in DMEM. In particular, the influence of 10 mM
20 carnosine, glycylphenylalanine (GlyPhe), Phe, or Gly supplementation of DMEM on protein content of MDCK cells was measured.

Figure 20 is a graph illustrating substrate (10mM) regulation of GlySar uptake by MDCK cells cultured in DMEM. In particular, the influence of 10 mM
25 carnosine, glycylphenylalanine (GlyPhe), Phe, or Gly supplementation of DMEM on H⁺-dependent uptake of [³H]Glycylsarcosine (GlySar) by MDCK cells was measured.

Figure 21 is a graph illustrating substrate (10mM) regulation of protein content of MDCK cells cultured in DMEM. In particular, the influence of 10 mM
glycylsarcosine (GlySar), glycylproline (GlyPro), glycylphenylalanine (GlyPhe), or carnosine of DMEM on protein content of MDCK cells was measured.

Figure 22 is a graph illustrating substrate (10mM) regulation of GlySar
30 uptake by MDCK cells cultured in DMEM. In particular, the influence of 10 mM

glycylsarcosine (GlySar), glycylproline (GlyPro), glycylphenylalanine (GlyPhe), or carnosine on H⁺-dependent uptake of [³H]Glycylsarcosine (GlySar) by MDCK cells was measured.

Figure 23 is a graph illustrating the influence of DMEM, nutrient depleted, dexamethasone (Dex), or insulin (ins) on H⁺-dependent uptake of [³H]Glycylsarcosine (GlySar) by MDCK cells.

Figure 24 is a graph illustrating influence of IGF-I on H⁺-dependent uptake of [³H]Glycylsarcosine (GlySar) by MDCK cells.

10

Definitions

The term "gene" is used broadly to refer to any segment of nucleic acid associated with a biological function. Thus, genes include coding sequences and/or the regulatory sequences required for their expression. For example, gene refers to a nucleic acid fragment that expresses mRNA, functional RNA, or specific protein, including regulatory sequences. Genes also include nonexpressed DNA segments that, for example, form recognition sequences for other proteins. Genes can be obtained from a variety of sources, including cloning from a source of interest or synthesizing from known or predicted sequence information, and may include sequences designed to have desired parameters.

20 The term "native gene" refers to gene that is present in the genome of an untransformed cell.

"Naturally occurring" is used to describe an object that can be found in nature as distinct from being artificially produced by man. For example, a protein or nucleotide sequence present in an organism (including a virus), which can be isolated from a source in nature and which has not been intentionally modified by man in the laboratory, is naturally occurring.

A "marker gene" encodes a selectable or screenable trait.

30 The term "chimeric gene" refers to any gene that contains 1) DNA sequences, including regulatory and coding sequences, that are not found together in nature, or 2) sequences encoding parts of proteins not naturally adjoined, or 3) parts

of promoters that are not naturally adjoined. Accordingly, a chimeric gene may comprise regulatory sequences and coding sequences that are derived from different sources, or comprise regulatory sequences and coding sequences derived from the same source, but arranged in a manner different from that found in nature.

5 A "transgene" refers to a gene that has been introduced into the genome by transformation and is stably maintained. Transgenes may include, for example, genes that are either heterologous or homologous to the genes of a particular cell to be transformed. Additionally, transgenes may comprise native genes inserted into a non-native organism, or chimeric genes. The term "endogenous gene" refers to a
10 native gene in its natural location in the genome of an organism. A "foreign" gene refers to a gene not normally found in the host organism but that is introduced by gene transfer.

 The terms "protein," "peptide" and "polypeptide" are used interchangeably herein.

15 Expression cassettes will comprise the transcriptional initiation region of the invention linked to a nucleotide sequence of interest. Such an expression cassette is provided with a plurality of restriction sites for insertion of the gene of interest to be under the transcriptional regulation of the regulatory regions. The expression cassette may additionally contain selectable marker genes.

20 The transcriptional cassette will include in the 5'-3' direction of transcription, a transcriptional and translational initiation region, a DNA sequence of interest, and a transcriptional and translational termination region. The termination region may be native with the transcriptional initiation region, may be native with the DNA sequence of interest, or may be derived from another source.

25 An oligonucleotide for use in probing or amplification reactions may be about 30 or fewer nucleotides in length (e.g., 9, 12, 15, 18, 20, 21 or 24, or any number between 9 and 30). Generally specific primers are upwards of 14 nucleotides in length. For optimum specificity and cost effectiveness, primers of 16-24 nucleotides in length may be preferred. Those skilled in the art are well
30 versed in the design of primers for use processes such as PCR. If required, probing

can be done with entire restriction fragments of the gene disclosed herein which may be 100's or even 1000's of nucleotides in length.

"Coding sequence" refers to a DNA or RNA sequence that codes for a specific amino acid sequence and excludes the non-coding sequences. It may constitute an "uninterrupted coding sequence", i.e., lacking an intron, such as in a cDNA or it may include one or more introns bounded by appropriate splice junctions. An "intron" is a sequence of RNA which is contained in the primary transcript but which is removed through cleavage and re-ligation of the RNA within the cell to create the mature mRNA that can be translated into a protein.

The terms "open reading frame" and "ORF" refer to the amino acid sequence encoded between translation initiation and termination codons of a coding sequence. The terms "initiation codon" and "termination codon" refer to a unit of three adjacent nucleotides ('codon') in a coding sequence that specifies initiation and chain termination, respectively, of protein synthesis (mRNA translation).

A "functional RNA" refers to an antisense RNA, ribozyme, or other RNA that is not translated.

The term "RNA transcript" refers to the product resulting from RNA polymerase catalyzed transcription of a DNA sequence. When the RNA transcript is a perfect complementary copy of the DNA sequence, it is referred to as the primary transcript or it may be a RNA sequence derived from post-transcriptional processing of the primary transcript and is referred to as the mature RNA. "Messenger RNA" (mRNA) refers to the RNA that is without introns and that can be translated into protein by the cell. "cDNA" refers to a single- or a double-stranded DNA that is complementary to and derived from mRNA.

"Regulatory sequences" and "suitable regulatory sequences" each refer to nucleotide sequences located upstream (5' non-coding sequences), within, or downstream (3' non-coding sequences) of a coding sequence, and which influence the transcription, RNA processing or stability, or translation of the associated coding sequence. Regulatory sequences include enhancers, promoters, translation leader sequences, introns, and polyadenylation signal sequences. They include natural and

synthetic sequences as well as sequences which may be a combination of synthetic and natural sequences. As is noted above, the term "suitable regulatory sequences" is not limited to promoters.

"5' non-coding sequence" refers to a nucleotide sequence located 5' (upstream) to the coding sequence. It is present in the fully processed mRNA upstream of the initiation codon and may affect processing of the primary transcript to mRNA, mRNA stability or translation efficiency. (Turner et al., Molecular Biotechnology, 3:225 (1995)).

"3' non-coding sequence" refers to nucleotide sequences located 3' (downstream) to a coding sequence and include polyadenylation signal sequences and other sequences encoding regulatory signals capable of affecting mRNA processing or gene expression. The polyadenylation signal is usually characterized by affecting the addition of polyadenylic acid tracts to the 3' end of the mRNA precursor.

The term "translation leader sequence" refers to that DNA sequence portion of a gene between the promoter and coding sequence that is transcribed into RNA and is present in the fully processed mRNA upstream (5') of the translation start codon. The translation leader sequence may affect processing of the primary transcript to mRNA, mRNA stability or translation efficiency.

The term "mature" protein refers to a post-translationally processed polypeptide without its signal peptide. "Precursor" protein refers to the primary product of translation of an mRNA. "Signal peptide" refers to the amino terminal extension of a polypeptide, which is translated in conjunction with the polypeptide forming a precursor peptide and which is required for its entrance into the secretory pathway. The term "signal sequence" refers to a nucleotide sequence that encodes the signal peptide.

The term "intracellular localization sequence" refers to a nucleotide sequence that encodes an intracellular targeting signal. An "intracellular targeting signal" is an amino acid sequence that is translated in conjunction with a protein and directs it to a particular sub-cellular compartment. "Endoplasmic reticulum (ER) stop transit

signal" refers to a carboxy-terminal extension of a polypeptide, which is translated in conjunction with the polypeptide and causes a protein that enters the secretory pathway to be retained in the ER. "ER stop transit sequence" refers to a nucleotide sequence that encodes the ER targeting signal.

- 5 "Promoter" refers to a nucleotide sequence, usually upstream (5') to its coding sequence, that controls the expression of the coding sequence by providing the recognition for RNA polymerase and other factors required for proper transcription. "Promoter" includes a minimal promoter that is a short DNA sequence comprised of a TATA- box and other sequences that serve to specify the site of
- 10 transcription initiation, to which regulatory elements are added for control of expression. "Promoter" also refers to a nucleotide sequence that includes a minimal promoter plus regulatory elements that is capable of controlling the expression of a coding sequence or functional RNA. This type of promoter sequence consists of proximal and more distal upstream elements, the latter elements often referred to as
- 15 enhancers. Accordingly, an "enhancer" is a DNA sequence which can stimulate promoter activity and may be an innate element of the promoter or a heterologous element inserted to enhance the level or tissue specificity of a promoter. It is capable of operating in both orientations (normal or flipped), and is capable of functioning even when moved either upstream or downstream from the promoter.
- 20 Both enhancers and other upstream promoter elements bind sequence-specific DNA-binding proteins that mediate their effects. Promoters may be derived in their entirety from a native gene, or be composed of different elements derived from different promoters found in nature, or even be comprised of synthetic DNA segments. A promoter may also contain DNA sequences that are involved in the
- 25 binding of protein factors which control the effectiveness of transcription initiation in response to physiological or developmental conditions.

The "initiation site" is the position surrounding the first nucleotide that is part of the transcribed sequence, which is also defined as position +1. With respect to this site all other sequences of the gene and its controlling regions are numbered.

- 30 Downstream sequences (i.e. further protein encoding sequences in the 3' direction)

are denominated positive, while upstream sequences (mostly of the controlling regions in the 5' direction) are denominated negative.

Promoter elements, particularly a TATA element, that are inactive or that have greatly reduced promoter activity in the absence of upstream activation are referred to as "minimal or core promoters." In the presence of a suitable transcription factor, the minimal promoter functions to permit transcription. A "minimal or core promoter" thus consists only of all basal elements needed for transcription initiation, e.g., a TATA box and/or an initiator.

"Inducible promoter" refers to those regulated promoters that can be turned on in one or more cell types by an external stimulus, such as a chemical, light, hormone, stress, or a pathogen.

"Operably-linked" refers to the association of nucleic acid sequences on single nucleic acid fragment so that the function of one is affected by the other. For example, a regulatory DNA sequence is said to be "operably linked to" or "associated with" a DNA sequence that codes for an RNA or a polypeptide if the two sequences are situated such that the regulatory DNA sequence affects expression of the coding DNA sequence (i.e., that the coding sequence or functional RNA is under the transcriptional control of the promoter). Coding sequences can be operably-linked to regulatory sequences in sense or antisense orientation.

"Expression" refers to the transcription and/or translation of an endogenous gene or a transgene in cells. For example, in the case of antisense constructs, expression may refer to the transcription of the antisense DNA only. In addition, expression refers to the transcription and stable accumulation of sense (mRNA) or functional RNA. Expression may also refer to the production of protein.

The analysis of transcription start points in practically all promoters has revealed that there is usually no single base at which transcription starts, but rather a more or less clustered set of initiation sites, each of which accounts for some start points of the mRNA. Since this distribution varies from promoter to promoter the sequences of the reporter mRNA in each of the populations would differ from each other. Since each mRNA species is more or less prone to degradation, no single

degradation rate can be expected for different reporter mRNAs. It has been shown for various eukaryotic promoter sequences that the sequence surrounding the initiation site ('initiator') plays an important role in determining the level of RNA expression directed by that specific promoter. This includes also part of the transcribed sequences. The direct fusion of promoter to reporter sequences would therefore lead to much suboptimal levels of transcription.

A commonly used procedure to analyze expression patterns and levels is through determination of the 'steady state' level of protein accumulation in a cell. Commonly used candidates for the reporter gene, known to those skilled in the art are 9-glucuronidase (GUS), growth hormone (GH), Chloramphenicol Acetyl Transferase (CAT) and proteins with fluorescent properties, such as Green Fluorescent Protein (GFP) from *Aequora victoria*. In principle, however, many more proteins are suitable for this purpose, provided the protein does not interfere with essential cell functions. For quantification and determination of localization a number of tools are suited. Detection systems can readily be created or are available which are based on e.g. immunochemical, enzymatic, fluorescent detection and quantification. Protein levels can be determined in cell extracts or in intact tissue using in situ analysis of protein expression.

Generally, individual transformed lines with one chimeric promoter reporter construct will vary in their levels of expression of the reporter gene. Also frequently observed is the phenomenon that such transformants do not express any detectable product (RNA or protein). The variability in expression is commonly ascribed to 'position effects, although the molecular mechanisms underlying this inactivity are usually not clear.

"Non-specific expression" refers to constitutive expression or low level, basal ('leaky') expression in undesired cells or tissues from a 'regulated promoter'.

"Antisense inhibition" refers to the production of antisense RNA transcripts capable of suppressing the expression of protein from an endogenous gene or a transgene.

"Co-suppression" and "transwitch" each refer to the production of sense RNA transcripts capable of suppressing the expression of identical or substantially similar transgene or endogenous genes (U.S. Patent No. 5,231,020).

"Homologous to" refers to the similarity between the nucleotide sequence of two nucleic acid molecules or between the amino acid sequences of two protein molecules. Estimates of such homology are provided by either DNA-DNA or DNA-RNA hybridization under conditions of stringency as is well understood by those skilled in the art (as described in Haines and Higgins (eds.), Nucleic Acid Hybridization, IRL Press, Oxford, U.K.), or by the comparison of sequence similarity between two nucleic acids or proteins.

The term "substantially similar" refers to nucleotide and amino acid sequences that represent equivalents of the instant inventive sequences. For example, altered nucleotide sequences which simply reflect the degeneracy of the genetic code but nonetheless encode amino acid sequences that are identical to the inventive amino acid sequences are substantially similar to the inventive sequences.

In addition, amino acid sequences that are substantially similar to the instant sequences are those wherein overall amino acid identity is 95% or greater to the instant sequences. Modifications to the instant invention that result in equivalent nucleotide or amino acid sequences is well within the routine skill in the art. Moreover, the skilled artisan recognizes that equivalent nucleotide sequences encompassed by this invention can also be defined by their ability to hybridize, under stringent conditions (0.1X SSC, 0.1% SDS, 65°C), with the nucleotide sequences that are within the literal scope of the instant claims.

"Transgene activation system" refers to the expression system comprised of an inactive transgene and a chimeric site-specific recombinase gene, functioning together, to effect transgene expression in a regulated manner. The specificity of the recombination will be determined by the specificity of regulated promoters as well as the use of wild-type or mutant site-specific sequences. Both elements of the system can be chromosomally integrated and inherited independently.

"Target gene" refers to a gene on the replicon that expresses the desired target coding sequence, functional RNA, or protein. The target gene is not essential for replicon replication. Additionally, target genes may comprise native non-viral genes inserted into a non-native organism, or chimeric genes, and will be under the control of suitable regulatory sequences. Thus, the regulatory sequences in the target gene may come from any source, including the virus.

"Transcription Stop Fragment" refers to nucleotide sequences that contain one or more regulatory signals, such as polyadenylation signal sequences, capable of terminating transcription. Examples include the 3' non-regulatory regions of genes encoding nopaline synthase and the small subunit of ribulose biphosphate carboxylase.

"Translation Stop Fragment" refers to nucleotide sequences that contain one or more regulatory signals, such as one or more termination codons in all three frames, capable of terminating translation. Insertion of a translation stop fragment adjacent to or near the initiation codon at the 5' end of the coding sequence will result in no translation or improper translation. Excision of the translation stop fragment by site-specific recombination will leave a site-specific sequence in the coding sequence that does not interfere with proper translation using the initiation codon.

"Blocking fragment" refers to a DNA fragment that is flanked by site specific sequences that can block the transcription and/or the proper translation of a coding sequence resulting in an inactive transgene. When the blocking fragment contains polyadenylation signal sequences and other sequences encoding regulatory signals capable of terminating transcription, it can block the transcription of a coding sequence when placed in the 5' non-translated region, i.e., between the transcription start site and the ORF. When inserted in the coding sequence a blocking fragment can block proper translation by disrupting its open reading frame. DNA rearrangement by site-specific recombination can restore transcription and/or proper translatability. For example, excision of the blocking fragment by site-specific recombination leaves behind a site-specific sequence that allows

transcription and/or proper translatability. A Transcription or Translational Stop Fragment will be considered a blocking fragment.

The terms "*in cis*" and "*in trans*" refer to the presence of DNA elements, such as the viral origin of replication and the replication protein(s) gene, on the same DNA molecule or on a different DNA molecule, respectively.

The terms "*cis*-acting sequence" and "*cis*-acting element" refer to DNA or RNA sequences whose functions require them to be on the same molecule. An example of a *cis*-acting sequence on the replicon is the viral replication origin.

The terms "*trans*-acting sequence" and "*trans*-acting element" refer to DNA or RNA sequences whose function does not require them to be on the same molecule.

"*Cis*-acting viral sequences" refers to viral sequences necessary for viral replication (such as the replication origin) and in *cis* orientation.

"Transactivating gene" refers to a gene encoding a transactivating protein. It can encode a viral replication protein(s) or a site-specific replicase. It can be a natural gene, for example, a viral replication gene, or a chimeric gene, for example, when regulatory sequences are operably-linked to the open reading frame of a site-specific recombinase or a viral replication protein. "Transactivating genes" may be chromosomally integrated or transiently expressed.

"Wild-type" refers to the normal gene, virus, or organism found in nature without any known mutation.

"Genome" refers to the complete genetic material of an organism.

The term "nucleic acid" refers to deoxyribonucleotides or ribonucleotides and polymers thereof in either single- or double-stranded form, composed of monomers (nucleotides) containing a sugar, phosphate and a base which is either a purine or pyrimidine. Unless specifically limited, the term encompasses nucleic acids containing known analogs of natural nucleotides which have similar binding properties as the reference nucleic acid and are metabolized in a manner similar to naturally occurring nucleotides. Unless otherwise indicated, a particular nucleic acid sequence also implicitly encompasses conservatively modified variants thereof

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(e.g., degenerate codon substitutions) and complementary sequences as well as the
sequence explicitly indicated. Specifically, degenerate codon substitutions may be
achieved by generating sequences in which the third position of one or more
selected (or all) codons is substituted with mixed-base and/or deoxyinosine residues
5 (Batzner et al., Nucleic Acid Res., 19, 5081 (1991); Ohtsuka et al., J. Biol. Chem.,
260, 2605 (1985); Rossolini et al., Mol. Cell. Probes, 8, 91 (1994)). A "nucleic acid
fragment" is a fraction of a given nucleic acid molecule. In higher animals,
deoxyribonucleic acid (DNA) is the genetic material while ribonucleic acid (RNA)
is involved in the transfer of information contained within DNA into proteins. A
10 "genome" is the entire body of genetic material contained in each cell of an
organism. The term "nucleotide sequence" refers to a polymer of DNA or RNA
which can be single- or double-stranded, optionally containing synthetic, non-
natural or altered nucleotide bases capable of incorporation into DNA or RNA
polymers. The terms "nucleic acid" or "nucleic acid sequence" may also be used
15 interchangeably with gene, cDNA, DNA and RNA encoded by a gene.

The invention encompasses isolated or substantially purified nucleic acid or
protein compositions. In the context of the present invention, an "isolated" or
"purified" DNA molecule or an "isolated" or "purified" polypeptide is a DNA
molecule or polypeptide that, by the hand of man, exists apart from its native
20 environment and is therefore not a product of nature. An isolated DNA molecule or
polypeptide may exist in a purified form or may exist in a non-native environment
such as, for example, a transgenic host cell. For example, an "isolated" or "purified"
nucleic acid molecule or protein, or biologically active portion thereof, is
substantially free of other cellular material, or culture medium when produced by
25 recombinant techniques, or substantially free of chemical precursors or other
chemicals when chemically synthesized. Preferably, an "isolated" nucleic acid is
free of sequences (preferably protein encoding sequences) that naturally flank the
nucleic acid (i.e., sequences located at the 5' and 3' ends of the nucleic acid) in the
genomic DNA of the organism from which the nucleic acid is derived. For
30 example, in various embodiments, the isolated nucleic acid molecule can contain

less than about 5 kb, 4 kb, 3 kb, 2 kb, 1 kb, 0.5 kb, or 0.1 kb of nucleotide sequences that naturally flank the nucleic acid molecule in genomic DNA of the cell from which the nucleic acid is derived. A protein that is substantially free of cellular material includes preparations of protein or polypeptide having less than about 30%,
5 20%, 10%, 5%, (by dry weight) of contaminating protein. When the protein of the invention, or biologically active portion thereof, is recombinantly produced, preferably culture medium represents less than about 30%, 20%, 10%, or 5% (by dry weight) of chemical precursors or non-protein-of-interest chemicals. Fragments and variants of the disclosed nucleotide sequences and proteins or partial-length
10 proteins encoded thereby are also encompassed by the present invention. By "fragment" is intended a portion of the nucleotide sequence or a portion of the amino acid sequence, and hence a portion of the polypeptide or protein, encoded thereby. Alternatively, fragments of a nucleotide sequence that are useful as hybridization probes generally do not encode fragment proteins retaining biological
15 activity. Thus, fragments of a nucleotide sequence may range from at least about 9 nucleotides, about 12 nucleotides, about 20 nucleotides, about 50 nucleotides, about 100 nucleotides or more.

By "variants" is intended substantially similar sequences. For nucleotide sequences, variants include those sequences that, because of the degeneracy of the
20 genetic code, encode the identical amino acid sequence of the native protein. Naturally occurring allelic variants such as these can be identified with the use of well-known molecular biology techniques, as, for example, with polymerase chain reaction (PCR) and hybridization techniques. Variant nucleotide sequences also include synthetically derived nucleotide sequences, such as those generated, for
25 example, by using site-directed mutagenesis which encode the native protein, as well as those that encode a polypeptide having amino acid substitutions. Generally, nucleotide sequence variants of the invention will have at least 40, 50, 60, to 70%, e.g., preferably 71%, 72%, 73%, 74%, 75%, 76%, 77%, 78%, to 79%, generally at least 80%, e.g., 81%-84%, at least 85%, e.g., 86%, 87%, 88%, 89%, 90%, 91%,

92%, 93%, 94%, 95%, 96%, 97%, to 98% sequence identity to the native nucleotide sequence.

By "variant" polypeptide is intended a polypeptide derived from the native protein by deletion (so-called truncation) or addition of one or more amino acids to the N-terminal and/or C-terminal end of the native protein; deletion or addition of one or more amino acids at one or more sites in the native protein; or substitution of one or more amino acids at one or more sites in the native protein. Such variants may results form, for example, genetic polymorphism or from human manipulation. Methods for such manipulations are generally known in the art.

Thus, the polypeptides of the invention may be altered in various ways including amino acid substitutions, deletions, truncations, and insertions. Methods for such manipulations are generally known in the art. For example, amino acid sequence variants of the polypeptides can be prepared by mutations in the DNA. Methods for mutagenesis and nucleotide sequence alterations are well known in the art. See, for example, Kunkel, Proc. Natl. Acad. Sci. USA, 82, 488 (1985); Kunkel et al., Methods in Enzymol., 154, 367 (1987); U. S. Patent No. 4,873,192; Walker and Gaastra, eds., Techniques in Molecular Biology, MacMillan Publishing Company, New York (1983) and the references cited therein. Guidance as to appropriate amino acid substitutions that do not affect biological activity of the protein of interest may be found in the model of Dayhoff et al., Atlas of Protein Sequence and Structure, Natl. Biomed. Res. Found., Washington, C.D. (1978), herein incorporated by reference. Conservative substitutions, such as exchanging one amino acid with another having similar properties, are preferred.

Thus, the genes and nucleotide sequences of the invention include both the naturally occurring sequences as well as mutant forms. Likewise, the polypeptides of the invention encompass both naturally occurring proteins as well as variations and modified forms thereof. Such variants will continue to possess the desired activity. The deletions, insertions, and substitutions of the polypeptide sequence encompassed herein are not expected to produce radical changes in the characteristics of the polypeptide. However, when it is difficult to predict the exact

effect of the substitution, deletion, or insertion in advance of doing so, one skilled in the art will appreciate that the effect will be evaluated by routine screening assays.

"Expression cassette" as used herein means a DNA sequence capable of directing expression of a particular nucleotide sequence in an appropriate host cell, comprising a promoter operably linked to the nucleotide sequence of interest which is operably linked to termination signals. It also typically comprises sequences required for proper translation of the nucleotide sequence. The coding region usually codes for a protein of interest but may also code for a functional RNA of interest, for example antisense RNA or a nontranslated RNA, in the sense or antisense direction. The expression cassette comprising the nucleotide sequence of interest may be chimeric, meaning that at least one of its components is heterologous with respect to at least one of its other components. The expression cassette may also be one which is naturally occurring but has been obtained in a recombinant form useful for heterologous expression. The expression of the nucleotide sequence in the expression cassette may be under the control of a constitutive promoter or of an inducible promoter which initiates transcription only when the host cell is exposed to some particular external stimulus. In the case of a multicellular organism, the promoter can also be specific to a particular tissue or organ or stage of development.

The proteins of the invention may be altered in various ways including amino acid substitutions, deletions, truncations, and insertions. Methods for such manipulations are generally known in the art. Methods for mutagenesis and nucleotide sequence alterations are well known in the art. See, for example, Kunkel, Proc. Natl. Acad. Sci. USA, 82:488-492 (1985); Kunkel et al., Methods in Enzymol. 154:367-382 (1987); US Patent No. 4,873,192; Walker and Gaastra, eds. (1983) Techniques in Molecular Biology (MacMillan Publishing Company, New York) and the references cited therein. Guidance as to appropriate amino acid substitutions that do not affect biological activity of the protein of interest may be found in the model of Dayhoff et al. (1978) Atlas of Protein Sequence and Structure (Natl. Biomed. Res. Found., Washington, D.C.), herein incorporated by reference.

Conservative substitutions, such as exchanging one amino acid with another having similar properties, may be preferred.

Thus, the genes and nucleotide sequences of the invention include both the naturally occurring sequences as well as mutant forms. Likewise, the proteins of the invention encompass both naturally occurring proteins as well as variations and modified forms thereof. Such variants will continue to possess the desired disease resistance activity. Obviously, the mutations that will be made in the DNA encoding the variant must not place the sequence out of reading frame and preferably will not create complementary regions that could produce secondary mRNA structure. See, EP Patent Application Publication No. 75,444.

The deletions, insertions, and substitutions of the protein sequence encompassed herein are not expected to produce radical changes in the characteristics of the protein. However, when it is difficult to predict the exact effect of the substitution, deletion, or insertion in advance of doing so, one skilled in the art will appreciate that the effect will be evaluated by routine screening assays. Hybridization of such sequences may be carried out under stringent conditions.

"Stringent hybridization conditions" and "stringent hybridization wash conditions" in the context of nucleic acid hybridization experiments such as Southern and Northern hybridization are sequence dependent, and are different under different environmental parameters. Longer sequences hybridize specifically at higher temperatures. An extensive guide to the hybridization of nucleic acids is found in Tijssen, Laboratory Techniques in Biochemistry and Molecular biology- Hybridization with Nucleic Acid Probes, page 1, chapter 2 "Overview of principles of hybridization and the strategy of nucleic acid probe assays" Elsevier, New York (1993). Generally, highly stringent hybridization and wash conditions are selected to be about 5° C lower than the thermal melting point (T_m) for the specific sequence at a defined ionic strength and pH. Typically, under "stringent conditions" a probe will hybridize to its target subsequence, but to no other sequences. For example, by "stringent conditions" or "stringent hybridization conditions" is intended conditions under which a probe will hybridize to its target sequence to a detectably greater

degree than to other sequences (e.g., at least 2- fold over background). By controlling the stringency of the hybridization and/or washing conditions, target sequences that are 100% complementary to the probe can be identified (homologous probing). Alternatively, stringency conditions can be adjusted to allow some mismatching in sequences so that lower degrees of similarity are detected (heterologous probing). Generally, a probe is less than about 1000 nucleotides in length, preferably less than 500 nucleotides in length.

Typically, stringent conditions will be those in which the salt concentration is less than about 1.5 M Na ion, typically about 0.01 to 1.0 M Na ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30° C for short probes (e.g., 10 to 50 nucleotides) and at least about 60° C for long probes (e.g., greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide.

Exemplary low stringency conditions include hybridization with a buffer solution of 30 to 35% formamide, 1 M NaCl, 1% SDS (sodium dodecyl sulphate) at 37° C, and a wash in 1X to 2X SSC (20X SSC = 3.0 M NaCl/0.3 M trisodium citrate) at 50 to 55° C. Exemplary moderate stringency conditions include hybridization in 40 to 45% formamide, 1.0 M NaCl, 1% SDS at 37° C, and a wash in 0.5X to 1X SSC at 55 to 60° C. Exemplary high stringency conditions include hybridization in 50% formamide, 1 M NaCl, 1% SDS at 37°C, and a wash in 0.1X SSC at 60 to 65° C.

Specificity is typically the function of post-hybridization washes, the critical factors being the ionic strength and temperature of the final wash solution. For DNA-DNA hybrids, the T_m can be approximated from the equation of Meinkoth and Wahl Anal. Biochem. 138:267-284 (1984); $T_m = 81.5^\circ \text{C} + 16.6 (\log M) + 0.41 (\% \text{GC}) - 0.61 (\% \text{form}) - 500/L$; where M is the molarity of monovalent cations, %GC is the percentage of guanosine and cytosine nucleotides in the DNA, % form is the percentage of formamide in the hybridization solution, and L is the length of the hybrid in base pairs. The T_m is the temperature (under defined ionic strength and

pH) at which 50% of a complementary target sequence hybridizes to a perfectly matched probe.

Very stringent conditions are selected to be equal to the T_m for a particular probe. An example of stringent hybridization conditions for hybridization of complementary nucleic acids which have more than 100 complementary residues on a filter in a Southern or Northern blot is 50% formamide with 1 mg of heparin at 42° C, with the hybridization being carried out overnight. An example of highly stringent conditions is 0.1 5 M NaCl at 72° C for about 15 minutes. An example of stringent wash conditions is a 0.2x SSC wash at 65° C for 15 minutes (see, Sambrook, *infra*, for a description of SSC buffer). Often, a high stringency wash is preceded by a low stringency wash to remove background probe signal. An example of medium stringency for a duplex of, e.g., more than 100 nucleotides, is 1x SSC at 45° C for 15 minutes. An example low stringency wash for a duplex of, e.g., more than 100 nucleotides, is 4-6x SSC at 40° C for 15 minutes. For short probes (e.g., about 10 to 50 nucleotides), stringent conditions typically involve salt concentrations of less than about 1.0M Na ion, typically about 0.01 to 1.0 M Na ion concentration (or other salts) at pH 7.0 to 8.3, and the temperature is typically at least about 30° C. Stringent conditions can also be achieved with the addition of destabilizing agents such as formamide. In general, a signal to noise ratio of 2x (or higher) than that observed for an unrelated probe in the particular hybridization assay indicates detection of a specific hybridization. Nucleic acids that do not hybridize to each other under stringent conditions are still substantially identical if the proteins that they encode are substantially identical. This occurs, e.g., when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code.

The following are examples of sets of hybridization/wash conditions that may be used to clone homologous nucleotide sequences that are substantially identical to reference nucleotide sequences of the present invention: a reference nucleotide sequence preferably hybridizes to the reference nucleotide sequence in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with

washing in 2X SSC, 0.1% SDS at 50°C, more desirably in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 1X SSC, 0.1% SDS at 50°C, more desirably still in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 0.5X SSC, 0.1% SDS at 50°C, preferably in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 0.1X SSC, 0.1% SDS at 50°C, more preferably in 7% sodium dodecyl sulfate (SDS), 0.5 M NaPO₄, 1 mM EDTA at 50°C with washing in 0.1X SSC, 0.1% SDS at 65°C.

T_m is reduced by about 1°C for each 1% of mismatching; thus, T_m , hybridization, and/or wash conditions can be adjusted to hybridize to sequences of the desired identity. For example, if sequences with >90% identity are sought, the T_m can be decreased 10°C. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (T_m) for the specific sequence and its complement at a defined ionic strength and pH. However, severely stringent conditions can utilize a hybridization and/or wash at 1, 2, 3, or 4°C lower than the thermal melting point (T_m); moderately stringent conditions can utilize a hybridization and/or wash at 6, 7, 8, 9, or 10°C lower than the thermal melting point (T_m); low stringency conditions can utilize a hybridization and/or wash at 11, 12, 13, 14, 15, or 20°C lower than the thermal melting point (T_m). Using these parameters, hybridization and wash compositions, and desired T , those of ordinary skill will understand that variations in the stringency of hybridization and/or wash solutions are inherently described. If the desired degree of mismatching results in a T of less than 45°C (aqueous solution) or 32°C (formamide solution), it is preferred to increase the SSC concentration so that a higher temperature can be used. An extensive guide to the hybridization of nucleic acids is found in Tijssen (1993) *Laboratory Techniques in Biochemistry and Molecular Biology-Hybridization with Nucleic Acid Probes*, Part 1, Chapter 2 (Elsevier, New York); and Ausubel et al., eds. (1995) *Current Protocols in Molecular Biology*, Chapter 2 (Greene Publishing and Wiley - Interscience, New York). See Sambrook et al. (1989) *Molecular*

Cloning: A Laboratory Manual (2d ed., Cold Spring Harbor Laboratory Press, Plainview, New York).

Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (T_m) for the specific sequence at a defined ionic strength and pH. However, stringent conditions encompass temperatures in the range of about 1°C to about 20°C, depending upon the desired degree of stringency as otherwise qualified herein. Nucleic acids that do not hybridize to each other under stringent conditions are still substantially identical if the polypeptides they encode are substantially identical. This may occur, e.g., when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code. One indication that two nucleic acid sequences are substantially identical is when the polypeptide encoded by the first nucleic acid is immunologically cross reactive with the polypeptide encoded by the second nucleic acid.

"Vector" is defined to include, inter alia, any plasmid, cosmid, or phage in double or single stranded linear or circular form which may or may not be self transmissible or mobilizable, and which can transform prokaryotic or eukaryotic host either by integration into the cellular genome or exist extrachromosomally (e.g. autonomous replicating plasmid with an origin of replication).

Specifically included are shuttle vectors by which is meant a DNA vehicle capable, naturally or by design, of replication in two different host organisms, which may be selected from actinomycetes and related species, bacteria and eucaryotic (e.g. higher cell, mammalian, yeast or fungal cells).

Preferably the nucleic acid in the vector is under the control of, and operably linked to, an appropriate promoter or other regulatory elements for transcription in a host cell such as a microbial, e.g. bacterial, or animal cell. The vector may be a bi-functional expression vector which functions in multiple hosts. In the case of genomic DNA, this may contain its own promoter or other regulatory elements and in the case of cDNA this may be under the control of an appropriate promoter or other regulatory elements for expression in the host cell.

"Cloning vectors" typically contain one or a small number of restriction endonuclease recognition sites at which foreign DNA sequences can be inserted in a determinable fashion without loss of essential biological function of the vector, as well as a marker gene that is suitable for use in the identification and selection of cells transformed with the cloning vector. Marker genes typically include genes that provide tetracycline resistance, hygromycin resistance or ampicillin resistance.

"Operably linked" means joined as part of the same nucleic acid molecule, suitably positioned and oriented for transcription to be initiated from the promoter. DNA operably linked to a promoter is "under transcriptional initiation regulation" of the promoter.

"Chimeric" is used to indicate that a DNA sequence, such as a vector or a gene, is comprised of more than one DNA sequences of distinct origin which are fused together by recombinant DNA techniques resulting in a DNA sequence, which does not occur naturally.

The terms "heterologous DNA sequence," "exogenous DNA segment" or "heterologous nucleic acid," as used herein, each refer to a sequence that originates from a source foreign to the particular host cell or, if from the same source, is modified from its original form. Thus, a heterologous gene in a host cell includes a gene that is endogenous to the particular host cell but has been modified through, for example, the use of DNA shuffling. The terms also include non-naturally occurring multiple copies of a naturally occurring DNA sequence. Thus, the terms refer to a DNA segment that is foreign or heterologous to the cell, or homologous to the cell but in a position within the host cell nucleic acid in which the element is not ordinarily found. Exogenous DNA segments are expressed to yield exogenous polypeptides.

A "homologous" DNA sequence is a DNA sequence that is naturally associated with a host cell into which it is introduced.

The following terms are used to describe the sequence relationships between two or more nucleic acids or polynucleotides: (a) "reference sequence",

(b) "comparison window", (c) "sequence identity", (d) "percentage of sequence identity", and (e) "substantial identity".

As used herein, "reference sequence" is a defined sequence used as a basis for sequence comparison. A reference sequence may be a subset or the entirety of a specified sequence; for example, as a segment of a full length cDNA or gene sequence, or the complete cDNA or gene sequence.

As used herein, "comparison window" makes reference to a contiguous and specified segment of a polynucleotide sequence, wherein the polynucleotide sequence in the comparison window may comprise additions or deletions (i.e., gaps) compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. Generally, the comparison window is at least 20 contiguous nucleotides in length, and optionally can be 30, 40, 50, 100, or longer. Those of skill in the art understand that to avoid a high similarity to a reference sequence due to inclusion of gaps in the polynucleotide sequence a gap penalty is typically introduced and is subtracted from the number of matches.

Methods of alignment of sequences for comparison are well known in the art. Thus, the determination of percent identity between any two sequences can be accomplished using a mathematical algorithm. Preferred, non-limiting examples of such mathematical algorithms are the algorithm of Myers and Miller, CABIOS 4:11-17 (1988); the local homology algorithm of Smith et al. Adv. Appl. Math. 2:482 (1981); the homology alignment algorithm of Needleman and Wunsch J. Mol. Biol. 48:443-453 (1970); the search-for-similarity-method of Pearson and Lipman Proc. Natl. Acad. Sci. 85:2444-2448 (1988); the algorithm of Karlin and Altschul, Proc. Natl. Acad. Sci. USA 87:2264 (1990), modified as in Karlin and Altschul, Proc. Natl. Acad. Sci. USA 90:5873-5877 (1993).

Computer implementations of these mathematical algorithms can be utilized for comparison of sequences to determine sequence identity. Such implementations include, but are not limited to: CLUSTAL in the PC/Gene program (available from Intelligenetics, Mountain View, California); the ALIGN program (Version 2.0) and GAP, BESTFIT, BLAST, FASTA, and TFASTA in the Wisconsin Genetics

Software Package, Version 8 (available from Genetics Computer Group (GCG), 575 Science Drive, Madison, Wisconsin, USA). Alignments using these programs can be performed using the default parameters. The CLUSTAL program is well described by Higgins et al. Gene 73:237-244 (1988); Higgins et al. CABIOS 5:151-153 (1989); Corpet et al. Nucleic Acids Res. 16:10881-90 (1988); Huang et al. CABIOS 8:155-65 (1992); and Pearson et al. Meth. Mol. Biol. 24:307-331 (1994). The ALIGN program is based on the algorithm of Myers and Miller, *supra*. The BLAST programs of Altschul et al., J. Mol. Biol. 215:403 (1990), are based on the algorithm of Karlin and Altschul *supra*. To obtain gapped alignments for comparison purposes, Gapped BLAST (in BLAST 2.0) can be utilized as described in Altschul et al. Nucleic Acids Res. 25:3389 (1997). Alternatively, PSI-BLAST (in BLAST 2.0) can be used to perform an iterated search that detects distant relationships between molecules. See Altschul et al., *supra*. When utilizing BLAST, Gapped BLAST, PSI-BLAST, the default parameters of the respective programs (e.g. BLASTN for nucleotide sequences, BLASTX for proteins) can be used. The BLASTN program (for nucleotide sequences) uses as defaults a wordlength (W) of 11, an expectation (E) of 10, a cutoff of 100, M=5, N=-4, and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength (W) of 3, an expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff & Henikoff, Proc. Natl. Acad. Sci. USA, 89, 10915 (1989)). See <http://www.ncbi.nlm.nih.gov>. Alignment may also be performed manually by inspection.

For purposes of the present invention, comparison of nucleotide sequences for determination of percent sequence identity disclosed herein is preferably made using the BlastN program (version 1.4.7 or later) with its default parameters or any equivalent program. By "equivalent program" is intended any sequence comparison program that, for any two sequences in question, generates an alignment having identical nucleotide or amino acid residue matches and an identical percent sequence identity when compared to the corresponding alignment generated by the preferred program.

As used herein, "sequence identity" or "identity" in the context of two nucleic acid or polypeptide sequences makes reference to the residues in the two sequences that are the same when aligned for maximum correspondence over a specified comparison window. When percentage of sequence identity is used in reference to proteins it is recognized that residue positions which are not identical often differ by conservative amino acid substitutions, where amino acid residues are substituted for other amino acid residues with similar chemical properties (e.g., charge or hydrophobicity) and therefore do not change the functional properties of the molecule. When sequences differ in conservative substitutions, the percent sequence identity may be adjusted upwards to correct for the conservative nature of the substitution. Sequences that differ by such conservative substitutions are said to have "sequence similarity" or "similarity." Means for making this adjustment are well known to those of skill in the art. Typically this involves scoring a conservative substitution as a partial rather than a full mismatch, thereby increasing the percentage sequence identity. Thus, for example, where an identical amino acid is given a score of 1 and a non-conservative substitution is given a score of zero, a conservative substitution is given a score between zero and 1. The scoring of conservative substitutions is calculated, e.g., as implemented in the program PC/GENE (Intelligenetics, Mountain View, California).

As used herein, "percentage of sequence identity" means the value determined by comparing two optimally aligned sequences over a comparison window, wherein the portion of the polynucleotide sequence in the comparison window may comprise additions or deletions (i.e., gaps) as compared to the reference sequence (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at which the identical nucleic acid base or amino acid residue occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the window of comparison, and multiplying the result by 100 to yield the percentage of sequence identity.

1 The term "substantial identity" of polynucleotide sequences means that a
polynucleotide comprises a sequence that has at least 70%, 71%, 72%, 73%, 74%,
75%, 76%, 77%, 78%, or 79%, preferably at least 80%, 81%, 82%, 83%, 84%, 85%,
86%, 87%, 88%, or 89%, more preferably at least 90%, 91%, 92%, 93%, or 94%,
5 and most preferably at least 95%, 96%, 97%, 98%, or 99% sequence identity,
compared to a reference sequence using one of the alignment programs described
using standard parameters. One of skill in the art will recognize that these values
can be appropriately adjusted to determine corresponding identity of proteins
encoded by two nucleotide sequences by taking into account codon degeneracy,
10 amino acid similarity, reading frame positioning, and the like. Substantial identity
of amino acid sequences for these purposes normally means sequence identity of at
least 70%, more preferably at least 80%, 90%, and most preferably at least 95%.

Another indication that nucleotide sequences are substantially identical is if
two molecules hybridize to each other under stringent conditions. Generally,
15 stringent conditions are selected to be about 5°C lower than the thermal melting
point (T_m) for the specific sequence at a defined ionic strength and pH. However,
stringent conditions encompass temperatures in the range of about 1°C to about
20°C, depending upon the desired degree of stringency as otherwise qualified herein.
Nucleic acids that do not hybridize to each other under stringent conditions are still
20 substantially identical if the polypeptides they encode are substantially identical.
This may occur, e.g., when a copy of a nucleic acid is created using the maximum
codon degeneracy permitted by the genetic code. One indication that two nucleic
acid sequences are substantially identical is when the polypeptide encoded by the
first nucleic acid is immunologically cross reactive with the polypeptide encoded by
25 the second nucleic acid.

The term "substantial identity" in the context of a peptide indicates that a
peptide comprises a sequence with at least 70%, 71%, 72%, 73%, 74%, 75%, 76%,
77%, 78%, or 79%, preferably 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%,
or 89%, more preferably at least 90%, 91%, 92%, 93%, or 94%, or even more
30 preferably, 95%, 96%, 97%, 98% or 99%, sequence identity to the reference

sequence over a specified comparison window. Preferably, optimal alignment is conducted using the homology alignment algorithm of Needleman and Wunsch J. Mol. Biol. 48:443-453 (1970). An indication that two peptide sequences are substantially identical is that one peptide is immunologically reactive with antibodies raised against the second peptide. Thus, a peptide is substantially identical to a second peptide, for example, where the two peptides differ only by a conservative substitution.

Detailed Description of the Invention

10 This invention relates to peptide amino acid absorption in the dog, and more particularly, to separate, whole or partial-length, complementary DNA encoding putative canine low-affinity, high-capacity H⁺/peptide transport proteins (cPepT1), mRNA transcripts corresponding to cPepT1, characterization of cPepT1 by glycylysarcosine (GlySar) uptake, identification of dipeptides, tripeptides, and tetrapeptides well recognized by cPepT1, and the effect of supplemental peptide
15 substrate on the transport capacity of cPepT1.

The invention also provides a pet food composition comprising at least one dipeptide, tripeptide, or tetrapeptide that provides enhanced uptake of amino acids by PepT1. A typical canine diet for use in the present invention may also, for
20 example, contain about 20 to about 30% crude protein, about 10 to about 20% fat, and about 10% total dietary fiber. However, no specific ratios or percentages of these or other nutrients are required.

The inventors have discovered a method for identifying peptides (e.g. dipeptides, tripeptides, or tetrapeptides) that increase transport of amino acids by PepT1 using MDCK cells, particularly when incubated with lactalbumin
25 hydrolysate and assayed at optimum time post-seeding, as indicated in Example 2.

In order that the invention may be more readily understood, reference is made to the following examples which are intended to illustrate the invention, but not limit the scope thereof.

30

EXAMPLE 1

Generation of Partial-length Canine PepT1 cDNA

Partial cloning of canine PepT1 (cPepT1) from small intestinal epithelium.

Initial attempts (over 150) to partially clone the putative canine PepT1
5 cDNA by reverse transcriptase-polymerase chain reaction (RT-PCR) methodology
failed. The source of mRNA was canine liver tissue that had been frozen for about 6
months (supplied by Dr. Randal Buddington, Mississippi State University) and
oligomer primers were based on the rabbit PepT1 sequence. Subsequently, frozen
canine "mid" small intestine (jejunal) tissue segments became available (supplied by
10 Dr. Buddington) and a partial length cDNA of about 780 base pairs (bp) was cloned
by RT-PCR. Total RNA was isolated from jejunal epithelium scraped from
intestinal sections using a standard acidic phenol-chloroform protocol. One μ g of
mRNA was isolated from total RNA using POLY A TRACT SYSTEM® (Promega,
Madison, WI) and reversed transcribed using murine leukemia virus reverse
15 transcriptase (Perkin Elmer, Foster City, CA) and oligo(dT) primers (Gibco BRL,
Grand Island, N.Y.). Successful PCR reactions were 50 μ L and contained 1 μ M
MgCl₂ and Taq polymerase (Perkin Elmer). Twenty-five thermal cycles of 94°C for
1 min, 40°C for 45 sec, and 72°C for 1 min were used. The cycles were preceded by
a 55 sec denaturization of the RT product at 95°C, followed by a 10 min extension
20 of RT-PCR products at 72°C. More than 150 RT-PCR reactions testing ten different
primer sets were required to achieve this protocol. The resulting cDNA using
Primer Set 4 (Figure 1) was TA-cloned into the pCR®II vector (Invitrogen, Carlsbad,
CA), plasmid-containing colonies selected by blue/white screening, and amplified
following instructions of the manufacturer. Restriction analyses of recovered
25 pCR®II/cDNA plasmids revealed that four of fifty-six clones contained cDNA
consistent with rabbit PepT1 cDNA (Figure 2).

Northern blot analysis of cPepT1 expression in dog tissue and MDCK cells.

The potential expression of cPepT1 mRNA by canine kidney, small
30 intestinal epithelium, and immortalized kidney distal tubule epithelial cells (Madin-

Darby Canine Kidney, MDCK) was evaluated by Northern analyses using cDNA derived from canine jejunal epithelium (Figure 3). RNA were subjected to 1% gel electrophoresis in the presence of 0.02 M formaldehyde, transferred by downward capillary action to 0.45- μ m nylon membranes (Hybond-N, Amersham, Arlington Heights, IL), and covalently cross-linked by ultra-violet light. cDNA were randomly labeled with [³²P]-CTP using a kit (Gibco BRL), purified through Sephadex-50 columns (Amersham Pharmacia, Piscataway, N.J.), and hybridized with blots at 56°C for 18 h. The blots were then washed 2 times at 56°C for 15 min and once at 57°C for 10 min. Autoradiographs were exposed to blots at 80°C for 24 h and the size of the transcript determined by regression of hybridized bands against the migration distance of 18S (1.9 kb) and 28S (4.9 kb) RNA.

Each canine tissue-derived cDNA (TA-clone 26, Figure 3A; TA-clone 6, Figure 3B) hybridized to three mRNA species in dog kidney, dog small intestinal epithelium, and MDCK cells. To confirm identification of PepT1 mRNA by these canine cDNAs, RNA isolated from dog kidney and liver tissues were probed for expression of PepT1 mRNA using a full-length rabbit PepT1 cDNA (Figure 4; rabbit PepT1 cDNA supplied by Drs. F. Leibach and V. Ganapathy, Medical College of Georgia). The results also demonstrated the expression of the same three PepT1 mRNA species by dog tissues, indicating that the full-length rabbit PepT1 cDNA and the cDNA derived from canine tissue in the present study identified the same transcripts. The mean/SD of transcript sizes calculated from these three blots were 4.2/.22, 2.75/.26, and 1.46/.42 kb, respectively. Collectively, these data indicate that liver, intestinal epithelial, and MDCK cells express the same size and number of PepT1 transcripts. In comparison, various tissues of chicken, sheep, cow, pig, rabbit, rat, human, and Caco2 cells are reported to express a single transcript, with the principle difference in size being between chicken (1.9) and mammalian species (2.8, 2.8, 2.9, 2.9, 3.0, 3.1, 2.9, respectively

Partial cloning and sequence identification of canine PepT1 (cPepT1) cDNA from MDCK cells.

To confirm the positive Northern analysis, identification of PepT1 mRNA expression using cDNA generated from dog small intestinal epithelium, RT-PCR methodologies were used to generate a PepT1 cDNA from MDCK cells. The target cDNA region was a subset of the cDNA generated by RT-PCR from canine small intestine (bp 83 to 887 of rabbit PepT1). Accordingly, PCR primers that corresponded to bp 259 to 619 of rabbit PepT1 (GenBank acc. no. U06467) were used to generate a partial-length "canine PepT1" (cPepT1) cDNA from mRNA isolated from MDCK cells. RNA was collected from cells that were plated at 30,000 cm² on rat tail collagen-coated dishes and cultured for 3 days in 10% fetal calf serum/DMEM. Reverse transcription of 5 µg of total RNA by SUPERScript® II reverse transcriptase (Gibco-BRL) was performed using random and oligo(dT) primers, per instructions of the manufacturer (Gibco-BRL). All PCR reactions contained 2 mM MgCl₂ and thermal cycling using Taq polymerase included 30 cycles at 94°C for 2 min, 55°C for 1 min, and 72°C for 2 min. The cycles were preceded by a 10 min denaturization of the RT product at 94°C, followed by a 10 min extension of RT-PCR products at 72°C. More than one hundred RT-PCR reactions were required to achieve this protocol.

The resulting cDNA of about 380 bp was TA-cloned, into the site of pCR®II vector (as described above), amplified, bacterial colonies evaluated by blue/white screening, and pCR®II/cDNA plasmids evaluated for cDNA by Eco RI/Pst I restriction analysis (as described above). Restriction analyses of recovered plasmids revealed that six of thirty-six clones contained cDNA consistent with rabbit PepT1 cDNA. Two of the confirmed plasmids were amplified in bacteria, recovered, and sent for sequencing by the University of Florida DNA Sequencing Core Facility (Gainesville). Sequence comparisons of this 380 bp cDNA (Figure 5) to PepT1 sequences of other species using BLAST 2.0.14. software (blast@ncbi.nlm.nih.gov) revealed that the canine sequence shares sequence homology of 79% to rabbit (bp 259 to 640; GenBank acc. no. 473375), 83% to rat (bp 213 to 593; GenBank acc. no.

D50664.1), 83% to mouse (bp 213 to 589; GenBank acc. no. AF205540), and 87% to human (bp 285 to 665; GenBank acc. no. 473375 and U13173) PepT1 sequences.

Demonstration of PepT1-like transport activity in MDCK cells.

5 As seen in Figures 3 and 5, MDCK cells express a canine homolog of
mammalian PepT1 mRNA. Potential expression of PepT1 transport activity (H^+ -
dependent, dipeptide inhibitable, low-affinity dipeptide transport) by confluent
MDCK cells was evaluated using whole-cell transport techniques and
glycylsarcosine (GlySar) as a model dipeptide substrate. Cells were seeded at
10 60,000 cells/cm² into 24-well trays that had been coated with rat tail collagen or
poly-L-lysine and cultured (95%O₂:5% CO₂ at 37°C) for 3 d in media consisting of
Dulbecco's Modified Eagle Medium/10% fetal calf serum/1% antimicrobial
antibacterial medium. Absorption (pmols/mg protein) of [³H]-glycyl-L-sarcosine
(GlySar; 6 mCi/mL, Moravsek Biochemicals, Brea, CA) was determined using the
15 24-well cluster tray method and representative scintillation counting. Before
transport, cells were incubated at 37°C for 30 min in 25 mM Hepes/Tris (pH 7.5),
140 mM NaCl, 5.4 mM KCl, 1.8 mM CaCl₂, 0.8 mM MgSO₄, and 5 mM glucose
(uptake buffer) to normalize intracellular amino acid and peptide pools. Transport
was initiated by the addition of 0.25 mL of uptake buffer that contained 2.88 μm
20 GlySar. After 30 min of uptake at 37°C, transport was terminated by rapid washing
of cells with 4 X 2 mL 4°C uptake buffer. Cellular protein was precipitated with
10% trichloroacetic acid and the supernatant recovered and counted to determine
radioactivity (³H) content. Cellular protein was then solubilized in 0.2 N NaOH and
0.2% SDS and quantified by the Lowry procedure, using bovine serum albumin as a
25 standard. The amount of H^+ -dependent GlySar absorbed was calculated as the
difference between uptake in pH 6.0 and pH 7.5 uptake buffers. The amount of
competitor substrate-inhibitable GlySar uptake was calculated as the quotient of
GlySar uptake in the absence and presence of 10 mM competitor substrate
(dipeptide or amino acid) multiplied by 100%.

GlySar uptake in the presence of an intracellularly H^+ gradient (extracellular pH of 6.0) was 2.3-fold higher in cells plated on collagen, and 1.7-fold higher when grown on poly-L-lysine, than uptake in pH 7.5 medium (Table 1). H^+ -dependent uptake of GlySar by MDCK cells was inhibited by 88 or 92% by the presence of 10 mM LeuTrp or TrpLeu when grown on collagen, and 87 or 92% when grown on poly-L-lysine, respectively (Table 1).

Table 1. Influence of extracellular pH and competitor substrates on uptake of [3H]-glycylsarcosine by MDCK cells cultured on collagen- or poly-L-lysine-coated trays. Cells were cultured as described in text and uptake compared in pH 7.5 or 6.0 media that contained 2.88 [3H]-glycylsarcosine for 30 min.

n	Extracellular pH	Competitor substrate(mM)	Glycylsarcosine uptake pmol 30 min ⁻¹ mg ⁻¹ protein	% inhibition of pH 6.0 uptake
Collagen-coated				
5	7.5	none	19.9 ± 2.80	na ¹
5	6.0	none	65.3 ± 7.95	100
5	6.0	LeuTrp (10)	7.68 ± 1.37	11.7
5	6.0	TrpLeu (10)	5.21 ± 0.39	8.0
4	6.0	Leucine (10)	63.0 ± 4.00	96.3
Poly-L-lysine-coated				
4	7.5	none	15.52 ± 1.06	na
5	6.0	none	42.31 ± 4.03	100
5	6.0	LeuTrp (10)	5.50 ± 0.58	13.0
5	6.0	TrpLeu (10)	3.44 ± 0.27	8.1
4	6.0	Leucine (10)	41.93 ± 2.70	100

¹ na, not applicable

To preliminarily characterize the kinetic parameters of peptide transport by MDCK cells, the uptake of GlySar in media that contained pH 6.0 and 0.00064, 0.0025, 0.010, 0.04, 0.160, 0.640, 2.56, or 10.2 mM of GlySar was measured (Figure 6). Total GlySar uptake was by a relatively low-affinity mechanism

(apparent K_m of about 4.0 mM) and high uptake velocity. Collectively, these characteristics of GlySar uptake are consistent with functional activity of PepT1 expressed by other species, as opposed to high-affinity, H^+ -dependent uptake by PepT2 (μM K_m). Accordingly, it is concluded that MDCK cells possess PepT1-like activity, consistent with detection of PepT1 mRNA by RT-PCR (Figures 1, 2, 5) and Northern blot analyses (Figures 3, 4).

Summary of Example 1

Separate partial-length canine PepT1 cDNAs (cPepT1) were generated by RT-PCR analyses from dog small intestinal epithelium ($n = 2$; Figures 1, 2) and immortalized canine kidney cells (MDCK cells, $n = 1$). The MDCK cDNA was sequenced (Figure 5) and found to share 79 to 87% sequence identity with PepT1 mRNA expressed by other mammalian species. Northern blot analyses using the intestinal epithelium-derived RT-PCR cDNA confirmed expression of canine PepT1 (cPepT1) by dog tissues (liver, $n = 3$; kidney, $n = 3$; small intestine $n = 1$) and MDCK cells ($n = 2$). The identification of mRNA transcripts corresponding to PepT1 using partial-length canine-derived PepT1 cDNA (Figure 3) was confirmed by hybridization to full-length rabbit cDNA (Figure 4). Characterization of GlySar uptake by MDCK cells demonstrated that MDCK cells express PepT1-like activity (Table 1, Figure 5), confirming detection of PepT1 mRNA expression by MDCK cells and use of MDCK cells as a model to characterize the function of canine PepT1.

EXAMPLE 2

Experimental Model of MDCK Cells for Evaluating the Effects of Various Peptide and Drug Substrates, and Hormones And/or Growth Factors, on the Expression of PepT1 Activity

Example 1 above showed that (1) a canine homolog of PepT1 (cPepT1) mRNA cloned from epithelia of the mid small intestine (jejunum) shares high sequence identity with PepT1 expressed by several other species, (2) canine liver,

kidney, and jejunal epithelium express a similar pattern of cPepT1 mRNA, and (3) MDCK cells are capable of H⁺-dependent peptide uptake. Accordingly, MDCK cells are an appropriate model to evaluate the biochemical characteristics of cPepT1. The specific goals of this research were to (1) characterize the functional activity of low-affinity H⁺-dependent GlySar uptake (PepT1 activity) by MDCK cells and (2) identify di- and tripeptides that are well recognized by cPepT1 (cPepT1), especially those that contain tryptophan and leucine.

Previous research (Brandsch et al., 1994, Biochem J. 299:253-260) briefly reported that H⁺-dependent peptide uptake by MDCK cells was greater when cells were grown in a medium that contained lactalbumin hydrolysate (LHM) versus one that contained free amino acids (DMEM). Therefore, in an attempt to establish the most sensitive model possible for evaluating peptide transport systems in MDCK cells, the potential influences of LHM (peptide-containing) versus DMEM (peptide-lacking) media, and subconfluent versus confluent initial cell plating densities were compared. MDCK cells were seeded at either 60,000 cells/well (subconfluent) or 120,000 cells/well (confluent) in DMEM and, after 1 d, cultured in DMEM or LHM media for 1, 2, 3, or 5 d. The amount of protein (index of cell growth) and GlySar uptake (index of peptide uptake capacity) expressed by each well of cells was then determined. As seen in Figure 7, the amount of cellular protein increased ($P < .05$) for both seeding densities and media with time of culture. A time x media interaction was observed, which reflects the greater protein content of cells grown in DMEM at day 6, as compared to those grown in LHM. At days 2, 3, or 4, however, no difference in protein content was observed.

The uptake of [³H]-GlySar (2.88 μM, 5 μCi/mL) by the MDCK cells described in Figure 7 was measured in the presence (pH 6.0 uptake buffer) and absence (pH 7.4 uptake buffer) of an extracellular-to-intercellular H⁺ (proton) gradient. A representative graph (Figure 8) compares the uptake of GlySar by cells seeded at 60,000/well and cultured in the LHM or DMEM. For both culture media, GlySar uptake in the presence of pH 6.0 was greater ($P < .01$) than that in pH 7.4 buffer and displayed a quadratic ($P < .01$) response to length of culture, reflecting a

buffer x day of culture interaction ($P < .01$). DMEM-cultured cells seeded at 120,000/well displayed almost identical uptake characteristics as just described for cells seeded at 60,000/well. In contrast, GlySar uptake in the presence of pH 6.0 buffer at day 3 by LHM-cultured cells was only 28% larger (quantitatively) than that observed by DMEM-cultured cells seeded at 60,000/well.

To further refine the analysis of media influence on the peptide transport capacity of MDCK cells plated at 60,000 or 120,000 cells per well, the H^+ -dependent GlySar uptake was calculated as the arithmetic difference between uptake in pH 6.0 and pH 7.4 buffers (Figure 9). Despite the comparable protein contents of cells observed at day 3 (Figure 7), cells seeded at 60,000 and grown in LHM media demonstrated about 60% greater capacity for GlySar uptake as did cells grown in DMEM (Figure 9; day x media interaction, $P < .01$). For all cells, the capacity for GlySar uptake per mg of cellular protein was decreased at day 6. This difference was the result of a lesser uptake at pH 6.0 by the LHM-cultured cells, and not the result of a larger pH 7.4 uptake.

The results of this experiment indicate that culturing cells in media that contains peptides does not increase growth rate but does increase the capacity for peptide uptake if cells are seeded at 60,000/well and cultured for 2 days in LHM. As such, these data are consistent with the induction of PepT1 expression by culture peptide-containing medium and describe an optimal set of culture conditions for characterizing H^+ -dependent peptide transport activity of the canine PepT1 transporter. These data also confirm, and more thoroughly describe, the stimulating effect of LHM versus DMEM media on peptide transport proteins that was initially reported by Brandsch et al. (1994).

Using the maximal uptake-stimulating culture parameters determined in Experiment 3, the effect of an extracellular-to-intracellular pH gradient on GlySar uptake was further evaluated to determine a pH level at which maximal GlySar uptake could be achieved, but which would replicate physiologic conditions (Figure 10). As expected, the presence of a pH gradient stimulated ($P < .001$) H^+ -dependent GlySar uptake, in a quadratic ($P < .01$) fashion. Uptake at pH 5.5 or 6.0

was about 2.7 times greater than that achieved at pH 7.5. These results are consistent with the data in Figures 8 and 9 and known H⁺-dependence of mammalian peptide transport proteins. Accordingly, the use pH 6.0 buffers for the characterization of H⁺-dependent GlySar uptake was incorporated into the standard experimental conditions.

To determine the appropriate time period to measure initial (linear) rates of GlySar uptake, a by-minute time-course experiment was performed. As seen in Figure 11, H⁺-dependent GlySar (100 uM) uptake increased linearly for 1 h and then slowed (quadratic response, $P < .01$). GlySar uptake in pH 6.0 buffer at 3.75, 7.5, 15, 30, 60 and 120 min was about 2, 2.1, 2.25, 2.65, 2.79, and 2.62 times more ($P < .001$), respectively, than uptake from pH 7.4 buffer. Because uptake was proportional to time of uptake through 1 h, future experiments were conducted using a 30-min time period.

To confirm that H⁺-dependent GlySar uptake was saturable, and therefore mediated, the uptake of GlySar from pH 6.0 and 7.4 uptake buffers containing 0.025, 0.1, 0.4, 1.6, 6.4, or 25.6 mM GlySar was evaluated (Figure 12). Uptake of GlySar was greatest ($P < .001$) from the pH 6.0 buffers, at all concentrations. H⁺-dependent GlySar uptake was saturable, consistent with an apparent K_m for GlySar of about 1.1 mM. These values are consistent with our preliminary trials that estimated a K_m of 1.1 mM for GlySar uptake by MDCK cells using only pH 6.0 uptake buffer and indicate that H⁺-dependent GlySar uptake is predominately, if not completely, a result of low affinity (mM) H⁺/peptide cotransporter activity (PepT1).

As a comparative value, the reported K_m of for GlySar uptake by the PepT1-expressing Caco-2 cells also is 1.1 mM. It is of interest also to note that GlySar uptake in the absence of a pH gradient (pH 7.4 buffers) also displayed linear ($P < .01$) and quadratic ($P < .001$) components, (1) reflects that the pH “7.4” buffer was in fact slightly acidic, (2) represents the activity of the putative basolateral peptide transporter running in “reverse”, or (3) indicates the presence of a non-characterized peptide transport system. As a result of this experiment, subsequent H⁺-dependent

peptide transport trials were conducted using 100 μ M GlySar, a value well below the K_m but one that will result in increased transport activity, and thus, sensitivity.

Characteristic hallmarks of low affinity H^+ /peptide cotransport activity, classically defined using membrane vesicles of several species, and more recently by functional expression studies using human, rat, and rabbit PepT1 cDNA, is the recognition of some, but not all, β -lactam antibiotics. In addition, PepT1 recognition of cefadroxil is low (the K_i of cefadroxil inhibition of GlySar uptake by PepT1 is 3 mM), whereas recognition of cefadroxil by PepT2 is high (the K_i of cefadroxil inhibition of PepT2 transport of GlySar is 30 μ M). To determine whether MDCK cPepT1 activity shared these functional features, the uptake of 100 μ M GlySar in the absence and presence of pH 7.5 and pH 6.0 buffer, and, in pH 6.0 buffers, the presence of 1 mM additional GlySar (self-inhibitor control), 3 mM Penicillin-G, 30 μ M cefadroxil, or 3 mM cefadroxil was compared (Figure 13). H^+ -dependent GlySar uptake was not inhibited by penicillin-G or 30 μ M cefadroxil, but was inhibited about 76% by 3 mM cefadroxil. As expected, the presence of 1 mM GlySar self-inhibited 100 μ M GlySar uptake by 64%. These results indicate that H^+ -dependent uptake of GlySar by MDCK cells is by PepT1 activity.

Other hallmarks of PepT1 function are the decreased ability of Gly-containing peptides to inhibit GlySar, in proportion to their length, and sensitivity to inhibition by carnosine (β -Ala-His). To determine if cPepT1 activity behaves as reported for other PepT1 activities, the relative abilities of 1 mM Gly ($[^3H]$ -Gly free amino acid control), GlyGly, $[Gly]_4$, or $[Gly]_5$ to inhibit H^+ -dependent 100 μ M GlySar was determined (Figure 14). Gly (5.0%) and $[Gly]_5$ (7.3%) did not influence uptake, whereas GlyGly inhibited and $[Gly]_4$ tended to inhibit uptake by 63 and 23%, respectively. This pattern of Gly-containing peptides to inhibit GlySar uptake in an inverse proportion to the number of glycyl residues in the canine MDCK cell model is consistent with PepT1 activities reported for other species. Similarly, GlySar uptake was inhibited 50% by 1 mM carnosine (data not shown but listed in Table 2 below).

Together with the molecular identification of PepT1 mRNA expression in MDCK cells using full-length rabbit cDNA and our canine RT-PCR product (*See Example 1 data*), the above biochemical characterization data indicate that H⁺-dependent GlySar uptake activity in MDCK cells is consistent with the low-affinity, high-capacity of the PepT1 transport protein. Collectively, the above experiments resulted in the generation of an experimental regimen for the culture and determination of H⁺-dependent peptide transport activity in MDCK cells, with which to evaluate the relative substrate preferences of canine PepT1 (cPepT1).

Accordingly, the following general regimen was used to perform a series of experiments that evaluated the relative abilities of candidate di- (primarily) and tri-peptides to inhibit GlySar uptake by endogenously expressed cPepT1 in MDCK cells:

1. Sixty thousand cells/well were plated into collagen-coated 24-well trays and cultured at 37°C in an atmosphere of 95% air/5% CO₂ in DMEM/10% FCS that contained antibiotics for 1 day.
2. The media was removed and cells were cultured in LHM/10% FCS/antibiotics for 1 day.
3. The media was removed and cells cultured in LHM/10% FCS (no antibiotics) for 20 h.
4. The media was removed and cells cultured for 30 min in air at 37°C in depletion medium (25 mM Hepes/Tris (pH 7.5), 140 mM NaCl, 5.4 mM KCl, 1.8 mM CaCl₂, 0.8 mM MgSO₄, and 5 mM glucose, to normalize intracellular nutrient pools.
5. Transport was initiated by replacing depletion medium with uptake medium (depletion medium adjusted to pH 6.0 or kept at pH 7.4) that contained 100 μM GlySar (at a specific activity of 5 μCi/mL, with [³H]-GlySar supplying 2.88% of total GlySar substrate) and (or) 1 mM of inhibiting peptide.

An inhibitory substrate concentration of 1 mM was selected because the literature indicates that typical K_m values for PepT1 ranges from 0.5 to 5 mM.

Therefore, by selecting an inhibitor concentration of 1 mM (not expected to completely inhibit uptake), our goal was to more finely delineate the relative abilities of candidate inhibitors than if the typical 5 mM inhibitor concentration (expected to achieve close to 100% inhibition of GlySar uptake) was used.

- 5 Candidate peptides were selected based on their containing Trp, Leu, Met, and (or) Arg, substrates. In total, 23 inhibitory peptides and 2 drug compounds were screened using this protocol.

- To determine the potential of Trp and Leu absorption as dipeptides by cPepT1, the ability of TrpLeu versus LeuTrp dipeptides to inhibit 100 μ M GlySar uptake was evaluated (Figure 15). The presence of either TrpLeu or LeuTrp in the pH 6.0 uptake buffer abolished H^+ -dependent GlySar uptake by 117% or 114%, respectively. In contrast, neither Leu nor Trp significantly influenced H^+ -dependent GlySar uptake. These results indicate that a lesser concentration of inhibitor would be required to delineate the relative recognition of TrpLeu and LeuTrp by cPepT1.
- 15 With regard to the mechanism of H^+ -independent GlySar uptake observed throughout these experiments, it is of interest to note that TrpLeu and LeuTrp inhibited H^+ -independent GlySar uptake by 36% and 46%, respectively.

- To further evaluate the potential of Trp to be absorbed in the form of peptides by cPepT1, the ability of TrpTrp, TrpGly, and TrpGlyGly to inhibit GlySar uptake was compared (Figure 16). As observed for TrpLeu (Figure 15), TrpTrp abolished H^+ -dependent GlySar uptake and inhibited H^+ -independent uptake by about 22%. TrpGly abolished H^+ -dependent GlySar uptake but did not influence H^+ -independent uptake. The tripeptide TrpGlyGly also significantly inhibited GlySar uptake, but to a lesser extent (73%) than did TrpTrp or TrpGly.
- 20

- 25 To determine the relative potential of other amino acids (Met, Arg, Lys, Phe, for example) to be absorbed in the peptide-bound form, additional GlySar competitive inhibition experiments were conducted using the above-described regimen and a variety candidate peptides at 1 mM. The results of these experiments are summarized in Table 2, which also includes those experiments described in
- 30 Figures 13, 14, 15, and 16 for comparative purposes.

Table 2. Influence of 1mM extracellular peptides and antibiotics on 100 μ M glycylsarcosine (GlySar) uptake¹ by MDCK cells.

5	Extracellular Substrate (1 mM)	% inhibition of H ⁺ -dependent GlySar uptake ⁴	n
	Positive control (model) substrates		
	GlyGly	89	8
10	[Gly] ₄	19	8
	[Gly] ₅	9	8
	Carnosine (β -AlaHis)	50	8
	Penicillin-G	0	8
	Cefadroxil ²	0	6
15	Cefadroxil ³	59	5
	Treatment substrates		
	<i>100% inhibition</i>		
	GlnGln	100	8
20	GlyLeu	115	8
	GlyMet	114	8
	LeuMet	114	8
	LeuTrp	113	8
	MetLeu	122	8
25	MetMet	100	8
	MetPhe	100	8
	MetPro	100	8
	TrpLeu	116	8
	TrpTrp	119	7
30	<i><100% inhibition</i>		
	GlnGlu	83	8
	MetGlu	93	8
	MetLys	88	8
	TrpGly	88	7
35	MetGlyMetMet (SEQ ID NO:10)	50	8
	TrpGlyGly	33	7
	LeuArg	32	8
	ArgLeu	32	8
40	¹ pmol mg ⁻¹ protein 30 min ⁻¹		
	² 30 μ M		
	³ 3 mM		

⁴That portion of the %inhibition value that exceeds 100, likely represents the ability to inhibit H⁺-independent GlySar uptake.

The inhibitors are listed within groupings in order of their relative ability to inhibit 100 μ M of GlySar uptake. In addition to the listed peptides, the constituent free amino acids were tested within the appropriate experiment to evaluate whether the peptide-bound or free amino acid was responsible for any affect on GlySar uptake. As expected, the presence of 1 mM constituent free amino acid did not influence GlySar uptake. Inhibition percentages of 50% indicate that the inhibitor substrate was recognized at least as well as was GlySar, given that the K_m of GlySar was determined to be about 1 mM (Figure 12) and that the substrate was present at 1 mM. Of the 19 treatment peptides evaluated, eleven abolished H^+ -dependent GlySar uptake, with seven of these also displaying the ability to inhibit H^+ -independent GlySar uptake. Of the remaining eight peptides tested, four displayed greater than 80% inhibition while four inhibited GlySar uptake by 50% or less. These results indicate that a wide variety of peptides of nutritionally important constituent amino acids are recognized by cPepT1.

Overall, the observation that cPepT1 activity was sensitive to a number of substrates is typical of PepT1 function. However, what was surprising was the large number of peptides that completely inhibited GlySar uptake. To establish a more sensitive relative inhibitory order among peptides that inhibited GlySar uptake by more than 80%, and, therefore, a more accurate potential for recognition, fourteen peptides were re-screened for their ability to inhibit 100 μ M GlySar uptake using the same cell culture and transport regimen but using only 10% of the previous inhibitor concentration (100 μ M). The data from an experiment to directly compare the ability of 100 μ M Trp-containing peptides are shown in Figure 17. All Trp-containing peptides inhibited H^+ -dependent GlySar uptake. However, TrpLeu inhibited more (92%) than did LeuTrp (58%), TrpTrp (62%), or TrpGly (45%). These values and the results of other experiments comparing the relative ability of Leu-, Met-, and Arg-containing peptides are listed in Table 3.

Table 3. Influence of 100 μ M extracellular peptides on 100 μ M glycylsarcosine (GlySar) uptake¹ by MDCK cells.

	Extracellular substrate (100 μ M) ²	% inhibition of H ⁺ -dependent GlySar uptake	n
5	Trp-containing peptides		
	TrpLeu	92	8
	TrpTrp	62	8
10	LeuTrp	58	8
	TrpGly	45	8
	Leu-containing peptides		
	TrpLeu	94	8
15	LeuMet	80	8
	MetLeu	77	8
	GlyLeu	65	8
	Met-containing peptides		
20	MetMet	85	8
	MetPhe	84	8
	MetGlu	31	8
	MetLys	30	8
25	Arg-containing peptides		
	ArgLeu	49	8
	LeuArg	8.9	8
	ArgTrp	8.9	8
30	¹ pmol mg ⁻¹ protein 30 min ⁻¹ ² Data are grouped by experiment		

Overall, four of the peptides inhibited GlySar uptake by at least 80%, six by more than 40%, and four less than 40%, thus establishing a relative ranking for recognition by cPepT1. Among the five Trp-containing peptides (Figure 17, Table 3), TrpLeu demonstrated the greatest ability to inhibit GlySar uptake. TrpLeu also demonstrated the greatest ability to inhibit GlySar uptake (94%) among the Leu-containing peptides. Among the Met-containing substrates that were directly compared within the same experiment, the neutral peptides, MetMet and MetPhe, inhibited more GlySar uptake than did the anionic (MetGlu) or cationic (MetLys)

carboxyl residues. Interestingly, as a group the Arg peptides demonstrated the least inhibitory ability, seemingly in keeping with the apparent lesser recognition by PepT1 of substrates with charged residues. However, it is of interest to note that 100 μ M ArgLeu demonstrated a much greater ability to inhibit GlySar uptake than did LeuArg (49 versus 8.9%).

To confirm the relative ranking of TrpLeu>TrpTrp inhibition of GlySar (Tables 2 and 3), Michaelis-Menton constants for substrate inhibition (K_i) of GlySar uptake by TrpLeu and TrpTrp were generated by graphical analyses of IC_{50} experiments (Figure 18). In keeping with the results achieved in the 100 μ M-inhibition studies, TrpLeu inhibited GlySar uptake at lower concentrations than did TrpTrp ($K_i = 0.2$ versus 0.75 μ M, respectively).

Collectively, the results of cPepT1 competitive inhibition trials using MDCK cells indicate that TrpLeu is better recognized by cPepT1 than any other tested peptide. The results also indicate that a number of Trp-, Leu, and Met-containing peptides also are well recognized by cPepT1. Ultimately, in the intestinal environment, it is the combination of recognition by the transporter and relative resistance of the peptide to luminal and membrane-bound peptidases that will determine how much of a given peptide will be absorbed. In this regard, there is some evidence to suggest that Gly-X peptides are more resistant than other peptides, especially by blood and renal peptidases. If so, then GlyLeu may be a better candidate substrate than TrpLeu to supply Leu. Similarly, tripeptides, as a group, are thought to be relatively resistant to hydrolysis. Thus, more TrpGlyGly may prove to be absorbed in larger amounts by the intestine than TrpLeu.

An important result of this set of experiments was the establishment of a sensitive experimental regimen/model to evaluate potential affecters of peptide transport capacity. Accordingly, this experimental model of MDCK cells grown in LHM affords an opportunity to evaluate the effects of various peptide and drug substrates, and hormones and (or) growth factors, on the expression of PepT1.

Thus, the culture of MDCK cells in LHM versus DMEM results in an increase of H^+ -dependent GlySar uptake ($K_m = 1.1$ mM) that is consistent with

mammalian PepT1-like activity. Using this stimulated model, the ability of twenty-three di- and tripeptides at 1 mM, and fourteen at 100 μ M, extracellular concentrations were screened for their ability to inhibit 100 μ M GlySar uptake, as an indicator of recognition by PepT1. Of the Trp- and (or) Leu-containing peptides evaluated, TrpLeu ($K_i = 0.2 \mu$ M) and LeuTrp ($K_i = 0.75 \mu$ M) demonstrated the greatest ability to inhibit GlySar uptake, with TrpLeu demonstrating a relatively higher affinity (lower K_i) for PepT1. Of the Met-containing peptides evaluated, four (MetMet, MetPhe, LeuMet, MetLeu) appear particularly well recognized by PepT1. In contrast, as a group, Arg-containing peptides displayed the least inhibition of PepT1 activity. Overall, these results indicate that cPepT1 is capable of recognizing a variety of di- and tripeptides, including, for example, those that contain leucine and tryptophan.

EXAMPLE 3

Experimental Model to Determine Whether the H^+ /peptide Transport Capacity Expressed by MDCK Cells Is Sensitive to Substrate Regulation

Trial 1:

Examples 1 and 2 above demonstrated that Madin-Darby canine kidney (MDCK) cells express PepT1 mRNA and characterized H^+ -dependent biochemical properties. Therefore, MDCK cells were chosen as the experimental model to determine whether the H^+ /peptide transport capacity expressed by MDCK cells is sensitive to substrate regulation. Research from Example 2 demonstrated that MDCK cells grown in lactalbumin hydrolysate medium (LHM) had elevated levels of peptide uptake capacity. Accordingly, to avoid potential confounding effects of the peptide-containing LHM and individual treatment peptides, DMEM (contains no peptides) and not LHM was selected as the appropriate medium to test the influence of extracellular peptides on canine PepT1 functional capacity of MDCK cells. GlyPhe was selected as a substrate because it has been reported to increase brush border membrane content of PepT1, (Shiraga T, Miyamoto K, Tanaka H, Yamamoto H, Taketani Y, Morita K, Tamai I, Tsuji A, Takada E. Cellular and

molecular mechanisms of dietary regulation on rat intestinal H⁺/peptide transporter PepT1. *Gastroenterology* 1999; 116:354-362), whereas Phe and Gly were tested as constituent free amino acid treatment controls. Carnosine was selected because of its high content in meat-based diets.

5 *Cell culture.* All cells were plated (60,000/2cm² well) and cultured (95% air/5% CO₂, 37°C) for 24 h in Dulbecco's Modified Eagle Media/10% fetal calf serum (FCS)/1%Antibiotic/ Antimicrobial solution (ABAM) (DMEM media). Following these initial common culture conditions, cells then were cultured in DMEM, or DMEM that contained 10 mM of Carnosine, GlyPhe, Phe, or Gly.

10 Media were changed every 24 h. Media treatments (n = 8) were as follows:

DMEM

DMEM + 10 mM Carnosine

DMEM + 10 mM GlyPhe

DMEM + 10 mM Phe

15 DMEM + 10 mM Gl

Uptake measurements. The measurement of [³H]Gly sarcosine uptake was performed by using a 24-well cluster tray method (Kilberg MS. Measurement of amino acid transport by hepatocytes in suspension and monolayer culture. *Methods Enzym* 1989; 173:564-575. Matthews JC, Aslanian A, McDonald KK, Yang W, Malandro MS, Novak DA, Kilberg MS. An expression system for mammalian amino acid transport using a stably maintained episomal vector. *Anal Biochem* 1997; 254:208-214), and used in Examples 1 and 2. Cells were cultured for 30 min in air at 37°C in depletion medium (25 mM Hepes/Tris (pH 7.5), 140 mM NaCl, 5.4 mM KCl, 1.8 mM CaCl₂, 0.8 mM MgSO₄, and 5 mM glucose), to normalize intracellular nutrient pools before transport. The transport assays are initiated by replacing depletion medium with uptake medium (Depletion medium adjusted to pH 6.0) that contained 100 μM GlySar (5 μCi/mL, with [³H]-GlySar supplying 2.88% of total GlySar). After a 30 minute incubation period, transport was terminated with four rinses of 4°C depletion medium (pH 7.5). Two hundred and twenty μL of 10% trichloroacetic acid was added to each well, and the radioactivity of the supernatant

quantified by liquid scintillation counting. The cells of each well are solubilized in 0.2 N NaOH/0.2% SDS and the protein quantified by using the modified Lowry assay, using bovine serum as a standard. *Id.* Peptide uptake will be reported as pmol* mg⁻¹ protein* 30 min⁻¹. Uptake measurements were taken after 24, 48, and 72 hours of culture in treatment media.

Results. The previous research characterizing H⁺-dependent peptide transport by MDCK cells (Example 2 above) clearly showed that transport velocity is dependent on protein content. Therefore, to make a valid comparison of various treatment parameters on GlySar uptake, the protein content of compared treatment groups must not differ. Accordingly, the influence of culture media on MDCK cellular protein was evaluated (Figure 19). All media treatments supported cellular growth from 1 to 3 d and no difference in protein content among treatments was observed. Similarly, no difference in uptake velocity (capacity) was observed among treatment groups, for any culture period (Figure 20).

Trial 2

The results from Trial 1 suggest that either canine PepT1 is not sensitive to substrate regulation or that the substrates and(or) stimulation time were inadequate to influence H⁺-dependent peptide uptake in MDCK cells. Again, DMEM was selected as the basal medium to allow the effect of individual peptides on peptide transport activity to be evaluated. To evaluate the latter two possibilities, a second trial was conducted that included a culture period of 9 d. GlySar was added as another potential affecter of H⁺-dependent peptide transport capacity because 10 mM GlySar it is reported capable of stimulating increased PepT1 activity (Adibi S. The oligopeptide transporter PepT1 in human intestine: biology and function. *Gastroenterology* 1997; 113:332-340) in Caco-2 cells. GlyPro was added as a treatment because of its high content in muscle tissue, thus is likely to be abundant in meat-based diets.

Cell culture. The MDCK cell line was maintained as described previously in the Methods section of Trial 1. Following initial and common culture conditions,

cells were cultured in DMEM, or DMEM that contained 10 mM GlySar, GlyPro, GlyPhe, or Carnosine. Media were changed every 24 h. Media treatments (n = 8) were as follows:

DMEM

- 5 DMEM + 10 mM GlySar
- DMEM + 10 mM GlyPro
- DMEM + 10 mM GlyPhe
- DMEM + 10 mM Carnosine

Uptake measurements. The measurement of [³H]GlySarcosine uptake was performed by using the 24-well cluster tray method as previously described in the Methods section of Trial 1. Peptide uptake will be reported as pmol* mg⁻¹ protein* 30 min⁻¹. Uptake measurements were taken after 4, 12, 24, 36, 72, 120, 168, and 216 hours of culture in treatment media.

Results. Protein content in all treatment groups increased linearly from 4 to 216 h (9 d) of culture, for all treatment groups (Figure 21). However, within a culture period, protein contents of treatment groups did not differ. Over the 216-h culture period, protein increased about 4.5 times, from about 40 to 220 µg/well. In contrast to Trial 1 results, media treatment did influence GlySar uptake capacity (Figure 22). In addition, a treatment x time effect was observed that represents differences in the time of culture required for GlySar and carnosine treatment stimulation of GlySar uptake capacity. Specifically, GlySar containing DMEM culture treatment resulted in an increase in GlySar uptake capacity of about 30% over DMEM control media by 24 h of culture time. This level of increase was maintained through 216 h. In contrast, culture in carnosine-containing media did not result in a significant (23%) increase of GlySar uptake capacity over that by DMEM-cultured cells until 72 h of culture. This stimulation then steadily increased to 291% over 216 h of culture. The nature of stimulated uptake between the two peptide substrates also differed. That is, the magnitude of carnosine-stimulated GlySar uptake was essentially constant from 72 to 216 h, whereas that for GlySar culture decreased during this period. Collectively, these data indicate that H⁺-

dependent peptide transport in cultured MDCK cells can be stimulated by at least two of PepT1 substrates, GlySar and carnosine.

Trial 3

5 The data from Trial 2 indicate that H⁺-dependent GlySar uptake capacity by fed MDCK cells can be upregulated by the inclusion of 10 mM GlySar for at least 24 h and 10 mM carnosine for at least 72 h. It is of equal interest to understand if H⁺-dependent GlySar uptake capacity is sensitive to nutrient deprivation and(or) stimulation by glucocorticoids. A preliminary study indicates that fasting increases
10 the expression of PepT1 in rat small intestine epithelia. Thamotharan M, Bawani S, Zhou X, Adibi S. Functional and molecular expression of intestinal oligopeptide transporter (PepT1) after a brief fast. *Metabolism* 1999; 48:681-684.

 To initiate investigation of potential influence of fasting and glucocorticoids on MDCK cells expression of GlySar uptake capacity, the H⁺-dependent uptake of
15 GlySar was evaluated over a 72 period of nutrient deprived or fed and cultured with dexamethasone (Dex) and compared to that by cells cultured in DMEM or DMEM that contained insulin (negative control) (Trial 3A). The “nutrient deprived” treatment actually contained 5 mM glucose and appropriate salts to ensure adequate basal metabolic conditions.

20 Although recruitment of PepT1 protein and activity appears sensitive to insulin-stimulated recruitment from cytosolic vesicles in Caco-2 cells (Thamotharan M, Bawani S, Zhou X, Adibi S. Hormonal regulation of oligopeptide transporter PepT1 after a brief fast. *Am J Physiol* 1999; 276:C821-826, MDCK cells are reported to be insensitive to insulin, likely as an inability to express the insulin
25 receptor. Hofmann C, Crettaz M, Bruns P, Hessel P, Hadawi G. Cellular responses elicited by insulin mimickers in cells lacking detectable plasma membrane insulin receptors. *J Cell Biol* 1985; 27:401-414. In contrast to the lack of insulin sensitivity, IGF-I is known to stimulate DNA synthesis and cell proliferation in MDCK cells. Sukegawa I, Hizuka N, Takano K, Asakawa K, Shizume K.
30 Characterization of IGF-1 receptors on MDCK cell line. *Endocrinol Japan* 1987;

34(3):339-346. Mouzon SH, Kahn R. Insulin-like growth factor-mediated phosphorylation and proto-oncogene induction in MDCK cells. *Mol Endocrinol* 1991; 5:51-60. The understanding that MDCK cells are apparently insensitive to insulin stimulation yet are sensitive to IGF-I stimulation appears to be a paradox given that the supraphysiologic levels of both substrates employed in the perspective studies and the known ability of insulin to cross react with the IGF-I receptor. Accordingly, another trial (Trial 3B) was conducted to evaluate the influence of increasing IGF-I concentrations on H⁺-dependent GlySar uptake by MDCK of the same plating stock.

10

Trial 3A:

Cell culture. MDCK cells were maintained as described in Trial 1, except that cells were cultured for only 1 d before transport trials were performed. Following initial and common culture conditions, cells were cultured in a “nutrient depleted” buffer (Hepes/Tris (pH 7.5), 140 mM NaCl, 5.4 mM KCl, 1.8 mM CaCl₂, 0.8 mM MgSO₄) that contained 5 mM glucose as an energy source, but that lacked amino acid or vitamin sources. In contrast, cells cultured in DMEM, or DMEM that contained 5 nM Dex, 500 nM Dex, 5 nM insulin, or 500 nM insulin, were adequately nourished. Media treatments (n = 4) were as follows:

20

Nutrient depleted

DMEM

DMEM + 5 nM Dex

DMEM + 500 nM Dex

DMEM + 5 nM Insulin

25

DMEM + 500 nM Insulin

Uptake measurements. The measurement of [³H]Glysarcosine uptake was performed by using the 24-well cluster tray method as previously described in the Methods section of Trial 1. Peptide uptake is reported as pmol* mg⁻¹ protein*30 min⁻¹. Uptake measurements were taken after 30 min and 4 h of culture in treatment media.

30

Trial 3B:

Trial 3B was conducted in the same manner as described for Trail 3A, except that cells were cultured in DMEM or DMEM that contained 1 nM IGF-1, 5 nM IGF-1, 25 nM IGF-1, or 100 nM IGF-1. Uptake measurements were taken after 30 min and 4 h of culture time. Media treatments (n = 4) were as follows:

DMEM (pH 6 measurement)

DMEM (pH 7.5 measurement)

DMEM + 1 nM IGF-1

DMEM + 5 nM IGF-1

DMEM + 25 nM IGF-1

DMEM + 100 nM IGF-1

Results. Protein content of the treatments within Trails 3A or 3B did not differ. After 4 h of culture, however, the capacity for H⁺-dependent peptide uptake was reduced 35% in cells deprived of nutrients but adequate in energy (Figure 23). In contrast, dexamethasone had no effect on GlySar uptake. As expected, and consistent with the concept that MDCK cells are insulin-insensitive, the presence of insulin for 4 h had no effect on GlySar uptake capacity. Similarly, culture of cells with increasing amounts of IGF-I elicited no significant stimulation of H⁺-dependent GlySar uptake (Figure 24). Quantitatively, however, 1 to 25 nM of IGF-I tended to increase GlySar uptake capacity by 10 to 15%.

Given the noted restrictions of Trail 3, and the low number of observations (n = 4) results from trial 3A and 3B suggest that H⁺-dependent uptake of GlySar by MDCK is sensitive to nutrient deprivation and, perhaps, IGF-I.

EXAMPLE 4

PepT1 Sequence

Clone12 (5th round; SEQ ID NO:11) Primer Pair is GSP3-4;GSP3-1R using regular RT-PCR

5 **catcttcttcacgtgg**tcaatgagttctgtgaaagatttctactatggaatgagagcactcctgattctgtacttcagacg
gttcacgggtgggacgataatctgtccacggccatctaccacacgtttgtggctctgtgctacctgacgccgatcctcgg
cgcactgatcgcagactcctggctgggaaagtcaagacaatcgtgtcactctccattgtctacacaattggacagggcgt
cactgcagtaagctcaattaatgacctcacagactataacaagatggaactcctgacaatctgtccgtgcatgtggcact
gtccatgattggcctggccctgatagctctgggaactggaggaataaagccctgtgtgtctgcatttggaggagaccagt
10 tgaagagggccaggaaaaacaagaacagattctttccatcttttatttggccattaatgctggaagcttgattccactat
tgtcactcccatgctcagagttcacgaatgtggaatttacagtcagaaagctgttaccactggcatttggggttcctgctg
ctctcatggcgtatctctgattgtattgtcattggcagtggaatgtacaagaagtttcagccccagggtaatgtcatgggt
aaagttgtcaagtgcattggtttgccctcaaaaataggttaggcaccggagtaagcagtttccaagaggaggactgg
ctggactgggctaaagagaaatacgaatgagcggctcatctctcaaatgaatggtcacaaaagtgtgtctgtacatcc
15 cactcccaatgttctgggccctgtttgaccagcagggtccagggtggacactgcaagcaacagctatgagtgaggaaaatt
ggacttctgaagttcagccagatcagatgcagactgtgaatgccatcttgattgtcgtcatgttccccatcatggatgccg
tggtgtacctctgattgcaaatgtggcttcaattcacctcctgaagaggatgacagttggaatgttctggcttccatgg
ccttcgtgatggcggcgattgttcagctggaaattgataaaactcttccagtcttcccaaaaacaaatgaagtccaaatcaa
agtactgaatataggaaatgggtccatgaatgtatctttctggagcgggtgacagttagccaaatgagtcataatcagat
20 ggatttatgacttttgatgtagacaaactgacaagtataaacatttcttccactggatcaccagtcattccagtactataact
ttgagcagggccatcgccatacccttctagtatgggcccccaataattaccgagtggtaaaggatggcctt**aaccagaa**
gccagaaaaaggag

Amplification conditions

25

	Initiale Denaturat	Denaturation	Annealing	Amplification	Extension	Cooling
Temp	94°C	94°C	55°C	72°C	72°C	4°C
Min.	10 min	2 min	1.5 min	2 min	10 min	inf.
Cycle	1		35		1	

Clone37 beginning (6th round; SEQ ID NO:12) Primer pair is GSP3-9; AUAP using 3'RACE Protocol

g c c a t c g c c a t a c c c t t c t a g t a t g g g c c c c a a t a a t t a c c g a g t g g t a a g g a t g g c c t t a a c c a g a a g c c a g a a a a
a g g a g a a a a t g g a a t c a g a t t t a a a t a g t c t t a a t g a g a g c c t c a a c a t c a c c a t g g g c g a c a a a g t t a t g t g a a t g t c
5 a c c a g t c a c a a t g c c a g c g a g t a t c a g t t c t t t t c t t t g g g c a c a a a a a c a t t a c a a a g t t c a a c a c a c a g a t c t c a c
a a a a t t g t a c a a a g t t c c a a t c a t c c a a c c t t g a a t t t g g t a g t g c a t a c c t a t g t a a t c g g a a c g c a g a g c a c t g g c
t g c c c t g a a t t g c a t a t g t t t g a a g a t a t t c a c c a a c a c a g t t a a c a t g g c t c t g c a g a t c c c g c a g t a c t t c c t c a t c a c c
t g c g g c g a g g t g g t t t c t c t g t c a c a g g a c t g g a g t t c a t a t t c a g g c c c c t c c a a c a t g a a g t c g g t g c t t c a g g
c g g g a t g g c t g c t g a c a g t g g c t t g t t g g c a a c a t c a t t g t g t c a t t g t g c a g g a g c a g g c c a g t t c a g t g a a c a g t g
10 g g c t g a a t a c a t c c t a t t t g c g g c a t t g c t t c t g g t t g t c t g t g t a a t a t t g c c a t c a t g g c c c g g t t t a c a c t t a c g t c a a t c
c a g c a g a g a t t g

Amplification conditions

	Initiale Denaturat	Denaturation	Annealing	Amplification	Extension	Cooling
Temp	94°C	94°C	52°C	72°C	72°C	4°C
Min.	10	2 min	1.5 min	2	10	inf.
Cycle	1		30		1	

15 Merge Sequence (SEQ ID NO:8) is:

c a t c t t c t c a t c g t g g t c a a t g a g t t c t g t g a a g a t t t c t a c t a t g g a a t g a g a g c a c t c c t g a t t c t g t a c t t c a g a c g g
t t c a t c g g g t g g g a c a t a a t c t g t c c a c g g c c a t c t a c c a c a c g t t t g t g g c t c t g t g t a c c t g a c g c c g a t c c t c g g c
g c a c t g a t c g c a g a c t c c t g g c t g g g a a g t t c a a g a c a a t c g t g t c a c t c t c a t t g t c t a c a a a t t g g a c a g g c g g t c
a c t g c a g t a a g t c a a t t a a t g a c c t c a c a g a c t a t a a c a a g a t g g a a c t c c t g a c a a t c t g t c c g t g c a t g t g g c a c t g t
20 c c a t g a t t g g c c t g g c c c t g a t a g c t c t g g g a a c t g g a g g a a a a g c c c t g t g t c t g c a t t t g g t g g a g a c c a g t t t g
a a g a g g g c c a g g a a a a c a a g a a c a g a t t c t t t c c a t c t t t a t t t g g c c a t a a t g c t g g a a g c t t g a t t c c a c t a t t g
t c a c t c c c a t g c t c a g a g t t c a c g a a t g t g g a a t t a c a g t c a g a a a g c t t g t a c c a c t g g c a t t t g g g t t c c t g c t g c t
c t c a t g g c c g t a t c t c t g a t t g t a t t t g c a t t g g c a g t g g a a t g t a c a a g a g t t c a g c c c a g g g t a a t g t a t g g g t a a
a g t t g t c a a g t g c a t t g g t t t t g c c c t c a a a a t a g g t t a g g c a c c g g a g t a a g c a g t t t c c a a g a g g g a g c a c t g g c t
25 g g a c t g g g c t a a a g a g a a t a c g a t g a g c g g c t a t c t c t c a a a t a a g a t g g t c a c a a a g t g a t g t t c t t g t a c a t c c c
a c t c c c a a t g t t c t g g g c c t g t t g a c c a g c a g g g c t c c a g g t g g a c a c t g c a a g c a a c a g c t a t g a g t g g g a a a a t t g
g a c t t c t t g a a g t t c a g c c a g a t c a g a t g c a g a c t g t g a a t g c c a t c t t a t t g t c g t a t g g t c c c c a t c a t g g a t g c c g t

ggtgtaccctctgattgcaaaatgtggcttcaattcacctccttgaagaggatgacagttggaatgttctggcttccatgg
 ccttcgtgatggcggcgattgttcagctggaaattgataaaactcttccagcttcccaaaacaaaatgaagtccaaatcaa
 agtactgaatataggaaatggtgccatgaatgtatcttttctggagcgggtggtgacagttagccaaatgagtcaatcagat
 ggatttatgacttttgatgtagacaaactgacaagtataaacatttcttccactggatcaccagtcattccagtgacttataact
 5 ttgagcagggccategccataacccttctagtatgggcccccaataattaccgagtggtaaaggatggccttaaccagaa
 gccagaaaaaggagaaaatggaatcagattataaatagtcttaatgagagcctcaacatcacatgggcgacaaagttt
 atgtgaatgtcaccagtcacaatgccagcgagtatcagttcttttcttgggcacaaaaaacattacaataagttcaacacaa
 cagatctcacaaaattgtacaaaagtctccaatcatccaaccttgaatttggtagtgcatafacctatgtaatcggaacgca
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 10 ttctcatcacctgcggcgaggtggtttctctgtcacaggactggagttctcatattctcaggccccctccaacatgaagtc
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acacttacgtcaatccagcagagattg

20709-204807

Multiple Alignment of Nucleotide Full Length Sequences

Sequence 1: XM_007063HomoSapiens 3045 bp
 Sequence 2: AY027496Ovis 2829 bp
 Sequence 3: D50306Rat 2900 bp
 Sequence 4: NM_053079Musmusculus 3128 bp
 Sequence 5: U13707Oryctolagusunic 2709 bp
 Sequence 6: AY029615Gallusgallus 2914 bp
 Sequence 7: SequencetosubmitGenbak 1840 bp

5

10 Start of Pairwise alignments

Aligning...

Sequences (4:5) Aligned. Score: 65
 Sequences (1:2) Aligned. Score: 65
 Sequences (2:3) Aligned. Score: 66
 Sequences (3:4) Aligned. Score: 88
 Sequences (4:6) Aligned. Score: 48
 Sequences (2:4) Aligned. Score: 64
 Sequences (1:3) Aligned. Score: 67
 Sequences (3:5) Aligned. Score: 66
 Sequences (4:7) Aligned. Score: 80
 Sequences (2:5) Aligned. Score: 77
 Sequences (3:6) Aligned. Score: 48
 Sequences (5:6) Aligned. Score: 51
 Sequences (1:4) Aligned. Score: 76
 Sequences (3:7) Aligned. Score: 81
 Sequences (5:7) Aligned. Score: 79
 Sequences (2:6) Aligned. Score: 50
 Sequences (6:7) Aligned. Score: 70
 Sequences (1:5) Aligned. Score: 67
 Sequences (2:7) Aligned. Score: 83
 Sequences (1:6) Aligned. Score: 49
 Sequences (1:7) Aligned. Score: 85

15

20

25

30

Guide tree file created:

[/net/nfs0/voll/production/w3nobody/tmp/999267.834538-239427.dnd]

Start of Multiple Alignment

There are 6 groups

Aligning...

5 Group 1: Sequences: 2 Score:48218
 Group 2: Sequences: 3 Score:43200
 Group 3: Sequences: 2 Score:42027
 Group 4: Sequences: 5 Score:39817
 Group 5: Sequences: 6 Score:30418
 Group 6: Sequences: 7 Score:33857

Alignment Score 249395

CLUSTAL-Alignment file created

[/net/nfs0/voll/production/w3nobody/tmp/999267.834538-239427.aln]

15 Your Multiple Sequence Alignment:

999267.834538-239427.aln

CLUSTAL W (1.81) multiple sequence alignment

20 D50306Rat (SEQ ID NO:3) -----CTGAACCTCCTGCTTG 15
 NM_053079Musmusculus -----
 (SEQ ID NO:4) -----
 XM_007063Homosapiens -----
 (SEQ ID NO:1) -----
 25 AY027496Ovis (SEQ ID NO:2) -----GAAACAACATCTTAGCACGGATTCTCCACCTGGACTCCTCGC 45
 U13707Oryctolagusunic -----
 (SEQ ID NO:5) -----
 SequencetosubmitGenbak -----
 (SEQ ID NO:7) -----
 30 AY029615Gallusgallus GCTCTGTCCGTCCCTCGGTCCCTCCGTCCGTCCGTCCCGCGGCGCG 50
 (SEQ ID NO:6) -----
 D50306Rat CCAGTCGCCGGTCAGGAGCCTCGGAGCCGCCACAAATGGGGATGTCCAAGT 65

5	NM_053079Musmusculus	---GTGCGCCGTCGCGAGCCCTTGGAGCCACCACAAATGGGATGTCCAAGT	47
	XM_007063Homo sapiens	-----GAATGTCCAAAT	12
	AY027496Ovis	TCGCCAGTCGCGAGGAGCCCTCGGAGCCGCCAGCATGGGAATGTCCGTGC	95
	U13707Oryctolagus cunic	-----CACCATGGGAATGTCTAAGT	20
	Sequencet osubmit Genbak	-----	
10	AY029615Gallus gallus	CCAGCAGCGTGC CGGCCCCCATGGCTGCAAAAAGTAAGAGTAAGGGCCGAT	100
	D50306Rat	CT---CGGGGTGCTTTGGCTACCCATTGAGCATCTTCTTCATCGTGGTC	112
	NM_053079Musmusculus	CT---CGGGGTGCTTCGGTTACCCGTTGAGCATCTTCTTCATCGTGGTC	94
	XM_007063Homo sapiens	CA---CACAGTTTCTTTGGTTATCCCTGAGCATCTTCTTCATCGTGGTC	59
	AY027496Ovis	CG---AAGAGTGTCTTCGGTTACCCCTTAAGCATCTTCTTCATCGTGGTC	142
15	U13707Oryctolagus cunic	CA---CTGAGTGTCTTCGGCTATCCCTGAGCATCTTCTTCATCGTGGTC	67
	Sequencet osubmit Genbak	-----CATCTTCTTCATCGTGGTC	19
	AY029615Gallus gallus	CAGTGCCGAAC TGCTTTGGCTACCCCTTGAGCATCTTCTTCATCGTGCATC	150
	D50306Rat	*****	**
	NM_053079Musmusculus	AATGAATTCGTGAAAGATTCTCCTACTATGGGATGCGAGCTCTCCTGGT	162
20	XM_007063Homo sapiens	AATGAATTCGTGAAAGATTCTCCTACTATGGCATGCGAGCACTCCTGGT	144
	AY027496Ovis	AATGAGTTTTCGCGAAAGATTTTCCCTACTATGGAATGCGAGCAATCCTGAT	109
	U13707Oryctolagus cunic	AATGAGTTCTGCGAAAGGTTCTCTTACTATGGAATGAGAGCACTCCTGAT	192
	Sequencet osubmit Genbak	AATGAGTTCTGCGAAAGGTTCTCCTACTATGGGATGAGAGCACTCCTGAT	117
	AY029615Gallus gallus	AATGAGTTCTGTGAAAGATTTTCCCTACTATGGAATGAGAGCACTCCTGAT	69
25	D50306Rat	AATGAGTTCTGCGAGAGGTTCTCCTACTATGGCATGCGAGCTGTGCTCGT	200
	NM_053079Musmusculus	*****	*
	XM_007063Homo sapiens	TCTGTACTTCAGGAACTTCCTTGGCTGGGATGATGACCTCTCCACGGCCA	212
	AY027496Ovis	TCTGTACTTCAGGAACTTCCTCGGCTGGGACGACAAATCTCTCCACGGCCA	194
	U13707Oryctolagus cunic	TCTGTACTTCACAAATTTTCATCAGCTGGGATGATAACCTGTCCACCGCCA	159
30	Sequencet osubmit Genbak	CCTGTACTTCCAAACGTTTCTCGGCTGGAACGACAACTGGGCACCGCCA	242
	AY029615Gallus gallus	TCTGTACTTCAGAAACTTCATCGGCTGGGACGACAACTGTCCACGGTCA	167
	D50306Rat	TCTGTACTTCAGACGTTTCATCGGCTGGGACGATAATCTGTCCACGGCCA	119
	NM_053079Musmusculus	ATTGTATTTCAAGTACTTCTCGGCTGGGATGACAACTTTTCTACAGCCA	250
	XM_007063Homo sapiens	*****	**
35	AY027496Ovis	*** * * * * * * * * * * * * * * *	*
	U13707Oryctolagus cunic	*** * * * * * * * * * * * * * * *	*
	Sequencet osubmit Genbak	*** * * * * * * * * * * * * * * *	*
	AY029615Gallus gallus	*** * * * * * * * * * * * * * * *	*
	D50306Rat	*** * * * * * * * * * * * * * * *	*

5	D50306Rat	GTGCGGAATCCACAGCCCAACAAGCTTGTACCCACTGGCCCTTGGGGTTC	662
	NM_053079Musmusculus	GTGCGGAATCCACAGTCAACAAGCTTGTACCCACTGGCCCTTGGGGTTC	644
	XM_007063Homo sapiens	ATGTGGAATTACAGTAAACAAGCTTGTACCCACTGGCCCTTGGGGTTC	609
	AY027496Ovis	ATGCGGAATTACAGTAAACAAGCTTGTACCCCTGGCCCTTGGGGTTC	692
	U13707Oryctolagus cunic	ATGTGGAATTACAGTAAACAAGCTTGTACCCACTGGCCCTTGGGATTC	617
10	Sequencet osubmit Genbak	ATGTGGAATTACAGTCAAGAAAGCTTGTACCCACTGGCATTTGGGGTTC	569
	AY029615Gallus gallus	ATGTGGCATTACAGCAGACAGCAGTGTCTACCCGCTGGCATTTGGAGTTC	700
		** ** ** *	
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		** ** ** *	
15	D50306Rat	CGGCAGCTCTCATGGCTGTGGCCCTAATTGTGTTTGTCTCTCGGCAGTGA	712
	NM_053079Musmusculus	CAGCGGCTCTCATGGCTGTGGCCCTAATTGTGTTTGTCTCTGGCAGTGA	694
	XM_007063Homo sapiens	CTGCTGCTCTCATGGCTGTAGCCCTGATTTGTGTTTGTCTCTGGCAGTGG	659
	AY027496Ovis	CTGCTGCACTCATGGCTGTATCTCTGATCGTGTGTTCATTCATTCGCAGTGA	742
	U13707Oryctolagus cunic	CTGCTATCTCATGGCTGTATCCCTGATCGTGTTCATCATCGGCAGTGGG	667
20	Sequencet osubmit Genbak	CTGCTGCTCTCATGGCCGTATCTCTGATTTGATTTGTTCATTCGCAGTGA	619
	AY029615Gallus gallus	CCGCTGCCCTCATGGCTGTTCATTTAGTTGTGTTCATAGCTGGAAGTGA	750
		* ** ***** ** *	
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		* ** ** *	
25	D50306Rat	ATGTACAAGAAGTTTCAGCCCCAGGGCAACATCATGGGCAAAAGTGGCCAA	762
	NM_053079Musmusculus	ATGTACAAGAAGTTCCAGCCCCAGGGCAACATCATGGGCAAAAGTGGCCAA	744
	XM_007063Homo sapiens	ATGTACAAGAAGTTCAAGCCACAGGGCAACATCATGGGTAAAGTGGCCAA	709
	AY027496Ovis	ATGTACAAGAAGTCCAGCCCCAGGGTAAACATCATGTCTAAAGTTGCCAG	792
	U13707Oryctolagus cunic	ATGTACAAGAAGTTCAAGCCCGAGGGGAACATCTGTAGCAAAAGTGGTGAA	717
30	Sequencet osubmit Genbak	ATGTACAAGAAGTTTCAGCCCCAGGGTAAATGTATGGGTAAAGTTGTCAA	669
	AY029615Gallus gallus	ATGTACAATAAAGTTTCAACCCGCAAGGCAATATAATGGTTTCGAGTTTGTAA	800
		***** ** * * * * *	
		* ** ** *	
		* ** ** *	
35	D50306Rat	GTGCATTGGCTTTGCCATCAAAAAACAGGTTTCGGCACCGAAGTAAGGCAT	812
	NM_053079Musmusculus	GTGCATTGGTTTGGCCATCAAAAAACAGGTTTCGGCACCGAAGTAAGGCAT	794
	XM_007063Homo sapiens	GTGCATCGGTTTGGCCATCAAAAAATAGATTTAGGCATCGGAGTAAGGCAT	759
	AY027496Ovis	GTGCATTGGTTTGGCCATCAAAAAATAGGATTAGCCATCGGAGTAAGAAAT	842
	U13707Oryctolagus cunic	GTGCATCTGTTTGGCCATCAAAAAATAGGTTTAGGCACCGCAGTAAGCAGT	767
40	Sequencet osubmit Genbak	GTGCATTGGTTTGGCCCTCAAAAAATAGGTTTAGGCACCGGAGTAAGCAGT	719
	AY029615Gallus gallus	ATGCATTGGATTGGCCATTAAAAACAGGTTTCGGCATCGCAGCAAAAGAT	850
		***** * ***** * ***** ** ** * * *	
		* ** ** *	
		* ** ** *	

D50306Rat	TTCCCAAGAGGGGAACACTGGCTGGACTGGCTAAAGAGAAAATACGATGAG	862
NM_053079Musmusculus	ATCCCAAGAGGGAGCACCTGGCTGGACTGGCTAAAGAGAAAATACGACGAG	844
XM_007063Homo sapiens	TTCCCAAGAGGGAGCACCTGGCTGGACTGGCTAAAGAGAAAATACGATGAG	809
AY027496Ovis	TTCTTAAGAGGGAGCACCTGGCTGGACTGGCTAGCGAGAAAATATGATGAG	892
U13707Oryctolagus cuniculus	TTCCCAAGAGGGGCGCACCTGGCTGGACTGGCTAAGAGAGAAAATACGACGAG	817
Sequencet osubmit Genbank	TTCCCAAGAGGGAGCACCTGGCTGGACTGGCTAAGAGAGAAAATACGATGAG	769
AY029615Gallus gallus	ATCCCAAAAGAGAGCACCTGGCTAGACTGGGCAAGCGAGAAAGTATGATAAA	900
	*** ** * * * ***** * ***** * ***** * * * * *	
D50306Rat	AGGCTCATCTCGCAGATTAAAGATGGTGACGAAGGTGATGTTCCCTGTACAT	912
NM_053079Musmusculus	CGGCTCATCTCACAGATTAAAGATGGTCACGAAGGTGATGTTCCCTGTTCAT	894
XM_007063Homo sapiens	CGGCTCATCTCCCAAATTAAAGATGGTTACGAGGGTGTGATGTTCCCTGTATAT	859
AY027496Ovis	CGGCTCATCTCTCAAATTAAAGATGGTTACAAAGGGTGTGATGTTCCCTGTACAT	942
U13707Oryctolagus cuniculus	CGGCTTATCGGCGCAGATCAAGATGGTTACGAGGGTGTGTTCCCTGTACAT	867
Sequencet osubmit Genbank	CGGCTCATCTCTCAAATTAAAGATGGTCACAAAAGTGTGTTCTTGTACAT	819
AY029615Gallus gallus	CGACTGATTGCTCAGACCACCAAGATGGTGTGAAGGTGCTTTTCCCTTTACAT	950
	* * * * * * * * * * ***** * * * * * * * * * *	
D50306Rat	TCCCTCCCCCATGTTTGGGCCCTTGTTTGACCAGCAGGGTTCAGGTGGA	962
NM_053079Musmusculus	CCCACTCCCCCATGTTTGGGCCCTTGTTTGACCACCAAGGGTCCAGATGGA	944
XM_007063Homo sapiens	TCCACTCCCAATGTTTGGGCCCTTGTTTGACCAGCAGGGCTCCAGGTGGA	909
AY027496Ovis	TCCTCTCCCCCATGTTTGGGCCCTTGTTTGATCAGCAGGGCTCCAGGTGGA	992
U13707Oryctolagus cuniculus	CCCACTCCCCCATGTTTGGGCCCTTGTTTGATCAGCAGGGTTCAGATGGA	917
Sequencet osubmit Genbank	CCCACTCCCAATGTTTGGGCCCTTGTTTGACCAGCAGGGCTCCAGGTGGA	869
AY029615Gallus gallus	CCCTCTCCCGATGTTTGGGCCACTTTTGTGACCAGCAGGGATCGAGATGGA	1000
	** ***** ***** * ***** * * * * * * * * * *	
D50306Rat	CACGTGCAAGCAACGACCACATGACTGGGAAAATTTGGAACAATTGAGATTTCAG	1012
NM_053079Musmusculus	CACGTGCAAGCAACGACCACATGAATGGGAAAATTTGAGCAAAATGAAAATTCAG	994
XM_007063Homo sapiens	CACGTGCAAGCAACCAACTATGTCGGGAAAATCGGAGCTCTTGAAAATTCAG	959
AY027496Ovis	CACGTGCAAGCAACGACCACATGATGGGAAAATTTGGAATCATTTGAAAATCCAG	1042
U13707Oryctolagus cuniculus	CGGTGCAAGCGACGACCACATGTCGGGAGAATTTGGAATCCTTGAAAATTCAG	967
Sequencet osubmit Genbank	CACGTGCAAGCAACAGCTATGATGGGAAAATTTGGAATCCTTGGAAGTTTCAG	919
AY029615Gallus gallus	CACGTGCAAGCCCAACTATGATGGGACTTTGGAGCTATGCAGATTTCAG	1050
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D50306Rat	CCGACCAGATGCAGACGGTGAACGCCCATCTTGATTGTCATCATGGTCCC	1062
NM_053079Musmusculus	CCGACCAGATGCAGACGGTGAATGCCATCCTGAATGTCAAATGGCCC	1044
XM_007063Homoapiens	CCCGATCAGATGCAGACCCGTGAACGCCATCCTGATCGTGATCATGGTCCC	1009
AY027496Ovis	CCGGATCAGATGCAGACGGTGAACGCCATCCTGATCGTCGTGATGGTCCC	1092
U13707Oryctolagusunic	CCGGATCAGATGCAGACTGTGAACACCATCTTGATTATTATCCTGGTCCC	1017
SequencetosubmitGenbak	CCAGATCAGATGCAGACTGTGAATGCCATCTTGATTGTGTCATGGTCCC	969
AY029615Gallusgallus	CCAGACCAAATGCAGACTGTCAATCCAATCCTGATTATAATAATGGTCCC	1100
	** ** * ***** ** ** * ** * ** * * ** *	
D50306Rat	CATTGTGGACGCCGTGGTGATCCCGTCAATGCAAAAATGTGGTTTCAACT	1112
NM_053079Musmusculus	CAATGTGGACGCCGTGTGTACCGTCAATGCAAAAATGTGGTTTCAACT	1094
XM_007063Homoapiens	GATCTTCGATGCTGTGTACCCCTCTCATGCAAAAATGTGGCTTCAATT	1059
AY027496Ovis	CATCGTGGATGCCGTGGTATATCCTCTGATCGCAAAAGTGTGGTTTAAATT	1142
U13707Oryctolagusunic	CATCATGGAACGCCGTGGTGATCCTCTGATTGCAAAAGTGTGGCCTCAACT	1067
SequencetosubmitGenbak	CATCATGGAATGCCGTGGTGATCCCTCTGATTGCAAAAATGTGGCTTCAATT	1019
AY029615Gallusgallus	AGTTGTAGATGCTGTGATTTATCCTTTAATCCAGAAAATGCAAGATCAATT	1150
	* ** * ** * ** * ** * ** * ** * ** * ** *	
D50306Rat	TCACCTCCCTGAAGAAGATGACCGTTGGGATGTTCTTGGCATCCATGGCC	1162
NM_053079Musmusculus	TCACATCCCTGAAGAAGATGACTGTTGGGATGTTCTTGGCGTCCATGGCC	1144
XM_007063Homoapiens	TCACCTCCTTGAAGAAGATGGCAGTTGGCATGGTCCTGGCCTCCATGGCC	1109
AY027496Ovis	TCACCTCCCTGAAGAAGATGACCGTCGGCATGTTTCTGGCCTCCATGGCT	1192
U13707Oryctolagusunic	TCACCTCTCTGAAGAAGATGACGATTGGGATGTTCTTGGCTTCCATGGCC	1117
SequencetosubmitGenbak	TCACCTCCTTGAAGAGGATGACAGTTGGAATGTTCTTGGCTTCCATGGCC	1069
AY029615Gallusgallus	TTACGCCCTTGAGGAGGATCACTGTTGGCATGTTCTTGGTGGTCTGGCT	1200
	* ** * ** * ** * ** * ** * ** * ** * ** *	
D50306Rat	TTTGTGGTGGCTGCAATTGTGCAGGTGGAATCGATAAAACTCTTCCAGT	1212
NM_053079Musmusculus	TTTGTGGTGGCTGCAATTGTGCAGGTGGAATCGATAAAACTCTTCCAGT	1194
XM_007063Homoapiens	TTTGTGGTGGCTGCCATCGTGCAGGTGGAATCGATAAAACTCTTCCAGT	1159
AY027496Ovis	TTCTGGCAGCTGCCATCGTGCAGGTGGACATGACAAAACTCTGCCCCGT	1242
U13707Oryctolagusunic	TTCTGGCAGCTGCAATCCTGCAGGTGGAATCGATAAAACTCTTCTCTGT	1167
SequencetosubmitGenbak	TTCTGTATGGCGCGATTTGTCAGCTGGAAATGATAAAACTCTTCCAGT	1119
AY029615Gallusgallus	TTCTGTGCTGCTGCTCTTTTGCAAGTGCAAAATAGATAAAACTCTTCCAGT	1250
	** ** * ** * ** * ** * ** * ** * ** * ** *	

5	D50306Rat	CTTCCCAGCGGAAATCAAGTTCAAAATTAAGGTCTTGAACATTTGGAAACA 1262
	NM_053079Musmusculus	CTTCCCTGGTGGAAATCAAGTCCAAATTAAGGTCTTGAACATCGGAAACA 1244
	XM_007063Homosapiens	CTTCCCAGGAAACGAAAGTCCAAATTAAGGTCTTGAATATAGGAAACA 1209
	AY027496Ovis	CTTCCCAGGAAATGAAGTCCAAATCAAAAGTCTTGAATATAGGAAATA 1292
	U13707Oryctolagusunic	CTTCCCAGGAAATGAAGTCCAAATTAAGGTCTTGAATGTAGGAAGTG 1217
10	SequencetosubmitGenbak	CTTCCCAGGAAATGAAGTCCAAATCAAAAGTCTTGAATATAGGAAATG 1169
	AY029615Gallusgallus	TTTCCCTGCAGCTGGACAGGCCCAAAATCAAAATAATAAATCTAGGTGATA 1300

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15	D50306Rat	ATGACATGGCCGTGTATTTTCCTGGAAAGAATGTGACAGTTGCCCAAAATG 1312
	NM_053079Musmusculus	ATAACATGACCGTGCATTTTCCTGGAAATAGTGTGACGCTTGCCCAAAATG 1294
	XM_007063Homosapiens	ATACCATGAATATATCTCTTCCTGGAGAGATGGTGACACTTGGCCCAATG 1259
	AY027496Ovis	ATAGCATGACCGTGTCTTTTCCCGGAAACGACAGTGCATGTGACCCAGATG 1342
	U13707Oryctolagusunic	AGAACATGATCATCTCTCTTCCTGGGACACGGTGACGCTCAACCCAGATG 1267
20	SequencetosubmitGenbak	GTGCCATGAATGTATCTTTTCCTGGAGCGGTGGTGACAGTTAGCCAAATG 1219
	AY029615Gallusgallus	GCAATGCGAATGT-TACATTTCTGCCCTAAATCTTCAGAACGTGACTGTCTT 1349
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25	D50306Rat	TCTCA--GACAGACACATT-CATGACTTTTCGATGTAGACCAGCTGACAA 1358
	NM_053079Musmusculus	TCTCA--GACAGACACGTT-CATGACTTTTCGATATAGACAAGCTGACAA 1340
	XM_007063Homosapiens	TCTCA--AACAAATGCATT-TATGACTTTTGTATGTAAACAAACTGACAA 1305
	AY027496Ovis	TCTCA--AACAAACGGATT-TCTGACTTTCAACGTAGACAACCT---AA 1385
	U13707Oryctolagusunic	TCTCA--AACGAATGAATT-CATGACTTTCAATGAAGACACACTGACAA 1313
30	SequencetosubmitGenbak	AGTCA--ATCAGATGGATT-TATGACTTTTGTATGTAGACAAACTGACAA 1265
	AY029615Gallusgallus	TCCCATGGAGTCAACAGGCTACAGGATGTTTGTAGTCTTCCAGCTAAAAAT 1399

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35	D50306Rat	GCATAAACGTGTCTTCTCCCGG-ATCTCCAGGCGTCACCACGGTAGCTCA 1407
	NM_053079Musmusculus	GCATAAACATATCTTCTCTTGG-ATCCCCAGGAGTCACCACAGTAGCTCA 1389
	XM_007063Homosapiens	GGATAAACATTTCTTCTCTCTGG-ATCACCAG--TCACTGCTGTAACTGA 1351
	AY027496Ovis	GTATAAACATTTCTTCTACTGG-AACACCAG--TCACTCCAGTAACTCA 1431
	U13707Oryctolagusunic	GCATAAACATCACTTCC--GG-ATCACAAG--TCACCATGATCACACC 1356
	SequencetosubmitGenbak	GTATAAACATTTCTTCCACTGG-ATCACCAG--TCATTCCAGTGACTTA 1311
	AY029615Gallusgallus	CTGTAATGGTAAATTTTGGGAGTGAGAGTAGAAGTGAAATAATATCGACTCA 1449

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[illegible]

D50306Rat	ATCAGTTTTTCCCTTCTGGCCAAAAGACTACACAATAAACACACCACAGA-	1653
NM_053079Musmusculus	ACAAGTTCCTCCCTTCTGGCGAAAAGCAGTACACAATAAACACACCAGGC-	1635
XM_007063Homosapiens	ACCAGTTTTTCCCTTCTGGCATAAAAGGCTTCAACAATAAGCTCAACAGA-	1597
AY027496Ovis	ATCTTTTTTCTCTTCTGGCGTAAAGAGCTTCAACAATAAACTCACCAGA-	1677
U13707Oryctolagusunic	ATCAGTTTTTCACTTCTGGAGTAAAGGGCTTCAACCGTCAGCTCGGCAGG-	1602
SequencetosubmitGenbak	ATCAGTCTTTTCTTGGGCACAAAACATTAACAATAAGTTCAACACAA	1558
AY029615Gallusgallus	ACAGTCCGTTCTCAGGAGGAACAATATATAGTATAGTATAACTGCAGG-	1697
	* * * * *	
D50306Rat	--GATTGCACCAAACTGTTCTATCTGATTTTAAATCTTCCAACCTTGACTT	1701
NM_053079Musmusculus	--GGTGGCACCAACCTGTCTAACTGATTTTAAATCTTCCAACCTTGACTT	1683
XM_007063Homosapiens	--GATTCCGCCACAATGTCAACCTAAATTTCAATACTTTCTACCTTGAATT	1645
AY027496Ovis	--GATTTACAAACAGTGTGAAAAACAGTTCAAAAATCTTACCTTGAATT	1725
U13707Oryctolagusunic	--CATCTCGGAGCAGTGCAGCGGGACTTTTGAGTCTCCGTACCTTGAAGTT	1650
SequencetosubmitGenbak	CAGATCTCACAAAATTTGTACAAAAGTTCTCCAATCATCCAACCTTGAATT	1608
AY029615Gallusgallus	-----TTCAACTAATTGCAAAACC-AACTTTCAGAG-----AAATTAGGATA	1736
	* * * * *	
D50306Rat	CGGCAGCGCGTACACCTACGTGATCAGAAAGTAGGGCGAGTGATGGCTGCC	1751
NM_053079Musmusculus	TGGCAGCGCGTATACCTACGTGATCCGA---AGGGCGAGTGATGGCTGCC	1730
XM_007063Homosapiens	TGGTAGTGCTTATACCTATATAGTCCAA---AGGAAGAATGACAGCTGCC	1692
AY027496Ovis	TGGTAGTGCGTTTACCTATGTAATCAGC---AGAAAAGAGTGACGGTTGCC	1772
U13707Oryctolagusunic	TGGCAGCGCGTACACGTACCTGATCAGC---AGCCAGGCTACTGGCTGCC	1697
SequencetosubmitGenbak	TGGTAGTGATATACCTATGTAATCGGA---ACGCAGAGCACCTGGCTGCC	1655
AY029615Gallusgallus	TGGTGGTGCTTATACGATCGTAATTAAAT---GAGTGTCTGGAGATGTGA	1783
	** * * * *	
D50306Rat	TGGAAGTGAAGGAATTCGAAGACATCCCAACCAACACCGGTGAACATGGCC	1801
NM_053079Musmusculus	TGGAAGTGAAGGAATTTGAAGACATCCCAACCAACACTGTGAACATGGCT	1780
XM_007063Homosapiens	CTGAAGTGAAGGTGTTTGAAGATATTTAGGCCAACACACAGTTAACATGGCT	1742
AY027496Ovis	CCGAACCAAGATTTTCGAAGACATCTCCCCCAACACAGTCAGCATGGCT	1822
U13707Oryctolagusunic	CCCAAGTGACGGAGTTTGAAGATATTTCCGCCCAACACATGAACATGGCT	1747
SequencetosubmitGenbak	CTGAATTGCATATGTTTGAAGATATTTCAACCAACACAGTTAACATGGCT	1705
AY029615Gallusgallus	CTCAATTAAGATACATTGAAGATATCCAAACCAATACAGTCCATATGGCT	1833
	** * * * *	

D50306Rat
 NM_053079Musmusculus
 XM_007063Homosapiens
 AY027496Ovis
 U13707Oryctolagusunic
 SequencetosubmitGenbak
 AY029615Gallusgallus

CTGCAGATCCACAGTACTTCTCCTCCTCACCTGCGGCGAGGTGGTCTTCTC 1851
 CTGCAGATCCACAGTACTTCTCCTTCTCACCTGCGGCGAGGTGGTCTTCTC 1830
 CTGCAAAATCCCGCAGTATTTCTTCTCACCTGTGGCGAAGTGGTCTTCTC 1792
 CTGCAGATCCCGCAGTACTTCTCCTCCTCACCTGTGGCGAGGTGGTCTTCTC 1872
 TGGCAAAATCCACAGTACTTCTCCTCATCACCTCTGGCGAGGTGGTCTTCTC 1797
 CTGCAGATCCCGCAGTACTTCTCCTCATCACCTGCGGCGAGGTGGTCTTCTC 1755
 TGGCAGATCCCTCAGTATTTTCATACTTACATGTGGAGAAGTAGTCTTCTC 1883
 *** ***** ** * * * * * ** ** ** **

5

D50306Rat
 NM_053079Musmusculus
 XM_007063Homosapiens
 AY027496Ovis
 U13707Oryctolagusunic
 SequencetosubmitGenbak
 AY029615Gallusgallus

TGTCACAGGACTGGAGTTCTCCTATTCCAGGCCCCGCTAACATGAAGT 1901
 TGTCACAGGACTGGAGTTCTCTTATTCCAGGCTCCGTCTAACATGAAGT 1880
 TGTACGGGATTGGAATTCTCATATTCTCAGGCTCCTTCCAAACATGAAGT 1842
 CATCACGGGCTGGAGTTCTCCTATTCTCAGGCTCCTTCCAAACATGAAGT 1922
 CATCACGGGCTGGAGTTCTCCTATTCTCAGGCTCCTTCCAAACATGAAGT 1847
 TGTCACAGGACTGGAGTTCTCATATTCTCAGGCCCCCTCCAAACATGAAGT 1805
 TGTCACTGGGCTGGAGTTTTCATACTACACAGGCACCATCTAATATGAAGT 1933
 **** ** ***** ** ** ** * * * * * ** ** **

15

D50306Rat
 NM_053079Musmusculus
 XM_007063Homosapiens
 AY027496Ovis
 U13707Oryctolagusunic
 SequencetosubmitGenbak
 AY029615Gallusgallus

CCGTGCTTCAGGCAGGATGGCTTCTAAACCGTGGCCATCGGTAATATCAT 1951
 CCGTGCTTCAGGCAGGCTGGCTTCTAACTGTGGCGGTTCGGCAATATCAT 1930
 CCGTGCTTCAGGCAGGATGGCTGCTGACCGTGGCTGTGGCAACATCAT 1892
 CCGTACTTCAGGCAGGATGGCTGTTGACCGTGGCCGTTGGCAACATCAT 1972
 CCGTGCTGCAGGACCGGTGGCTGCTGACCGTGGCTGTGGCAACATCAT 1897
 CCGTGCTTCAGGCGGGATGGCTGCTGACAGTGGCT----- 1840
 CAGTGCTGCAAGCAGGATGGCTGCTAACAGTGGCTGTCGGTAACATAATT 1983
 * ** ** * * ***** * ** *****

25

D50306Rat
 NM_053079Musmusculus
 XM_007063Homosapiens
 AY027496Ovis
 U13707Oryctolagusunic
 SequencetosubmitGenbak
 AY029615Gallusgallus

GTCTCATTTGTGGCTGAGGCAGGCCACTTCGACAAACAGTGGGTGAGTA 2001
 GTGCTCATCGTGGCAGGGCGGGGGCACTTCCCAAAACAGTGGGTGAGTA 1980
 GTGCTCATCGTGGCAGGGCGAGGCCAGTTCAGCAAAACAGTGGGCCGAGTA 1942
 GTGCTTATTGTGGCAGGAGCAGGCCAGTTCAGTGAACAGTGGGCCGAGTA 2022
 GTGCTCATCGTGGCGGGCGGGCCAGATCAACAGCAGTGGGCCGAGTA 1947

 GTCTTATTCGTGGCTGGAGCATCCAAACTCAGTGAGCAGTGGGCAGATA 2033

35

	D50306Rat	TGTTCTGTTCCGCTCCTTGCTCTCGTTCGTGCATCATATTGCCATT	2051
	NM_053079Musmusculus	CATTCTGTTTGCCCTCATTTGCTTCTGGTTGTTCTGCGTGATATTGCGCCATCA	2030
	XM_007063Homosapiens	CATTCTATTGCGCGTTGCTTCTGGTCTGTGTGTAATTTTGGCCATCA	1992
	AY027496Ovis	CGTTCTGTTTGGCGCATTTGCTTCTGGTCTGTGCATAAATATTGCGCCATCA	2072
5	U13707Oryctolagusunic	CATCCTCTTTGCGGCCCTGCTCTCTGGTCTGTGTCTCATATTGCGCCATCA	1997
	SequencetosubmitGenbak	-----	
	AY029615Gallusgallus	TGTTCTCTTTTGCTGCCTTGCTTTTTTGACAGTTTGCAATTATTTTGCTGTCA	2083
10	D50306Rat	TGGCCCGGATTCTACACCTACATCAACCCAGCACAGATCGAGGCACAGTTTC	2101
	NM_053079Musmusculus	TGGCTCGATTCTACACCTACATCAACCCAGCACAGAGATTGAAGCACAGTTT	2080
	XM_007063Homosapiens	TGGTCGGTCTTACTATTACATCAACCCAGCGGAGATCGAAGCTCAATTT	2042
	AY027496Ovis	TGGTCGATTCTATACGTATGTCAACCCCGCAGAGATTGAAGCTCAGTTT	2122
15	U13707Oryctolagusunic	TGGTCGATTCTATACGTATGTCAACCCGGCCGAGATCGAGGCTCAGTTT	2047
	SequencetosubmitGenbak	-----	
	AY029615Gallusgallus	TGGCATATTTTTTATACATATACTGATCCAATGAGGTTGAAGCCCCAACTT	2133
20	D50306Rat	GATGAGGATGAGAAGAAAAAGGGCGTAGGGAAGGAA--AACCCGTATTTC	2148
	NM_053079Musmusculus	GATGAGGATGAGAAGAAAAAGGGCATAGGAAAAGGAA--AACCCGTATTTC	2127
	XM_007063Homosapiens	GATGAGGATGAAAAGAAAAACAGACTGGAAAAGAGT---AACCCATATTTT	2089
	AY027496Ovis	GATGAGGATGACAAGGAGGATGACCTGGAAAAGAGT---AACCCATACGC	2169
	U13707Oryctolagusunic	GAAGAAAGATGAGAAGAAAAAGAACCCAGAAAAAGAAC--GACCTCTACCC	2094
	SequencetosubmitGenbak	-----	
25	AY029615Gallusgallus	GATGAAGAAGAAAAAGAAACAAATAAAAAACAGGATCCAGACTTGACACGG	2183
30	D50306Rat	CTCG----TTGGAACCTGTCTCTCAGACACAAACATGTGAAGATCAGAAAAGCA	2195
	NM_053079Musmusculus	TTCA----TTGGAACCAGTCTCACAGACAAATATGTGAAGGGCAGAAGGCA	2174
	XM_007063Homosapiens	CATG----TCAGGGGCCAATTTCAGAAAAACAGATGTGAAGGTCAGGAGGCA	2136
	AY027496Ovis	CAAG----CTGGACTTCGTCTCACAGACACAAATGTGAATGTCAAGGAAGCA	2216
	U13707Oryctolagusunic	CTCC----GTGGCGCCCCTCTCTCACAGACACAGATGTGA--GTCTGGAGGCG	2139
	SequencetosubmitGenbak	-----	
	AY029615Gallusgallus	AAAAAGAACTGAAGCTGTCTCTCAGATGTAGAAG-GTGTATTTCAAGAGCA	2232
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D50306Rat	-----
NM_053079Musmusculus	TTTTAAATTTTTTTTATTTTTTTTATTTTTTTTGGCTGTTTGTTCGA 2420
XM_007063Homosapiens	GTCCAGGCTGGAGTGCAATGGCACGATCTCAGCTCACTGC-----A 2364
AY027496Ovis	-----
U13707Oryctolagusunic	-----
SequencetosubmitGenbak	-----
AY029615Gallusgallus	-----
5	
D50306Rat	-----
NM_053079Musmusculus	GACAGGGTTTCTCGTGTGTAGCCCTTGGTGTCTGGAACTCACTCTGTA 2470
XM_007063Homosapiens	ACCTCCGCCCTCCAGGTTCAAGTAATTCTCCTGCCCTCAGCCTCCCGAGTG 2414
AY027496Ovis	-----
U13707Oryctolagusunic	-----
SequencetosubmitGenbak	-----
AY029615Gallusgallus	-----
10	
D50306Rat	-----
NM_053079Musmusculus	GACCAGACTGGCCTCAAACTCAGAAATCCACCTGCCCCCTGCCCTGCCCC 2520
XM_007063Homosapiens	GCTGGGATTAGCGGCA-----
AY027496Ovis	-----
U13707Oryctolagusunic	-----
SequencetosubmitGenbak	-----
AY029615Gallusgallus	-----
20	
D50306Rat	-----
NM_053079Musmusculus	TGCCCCCTGCCCCCTGCCCTCTGCCTCTGCCCTCCCAAGTGTGGATT 2570
XM_007063Homosapiens	-----TGCACCAACCACGCCCGAGCTATTTTGTATTTTAGTAGAGAT-- 2472
AY027496Ovis	-----
U13707Oryctolagusunic	-----
SequencetosubmitGenbak	-----
AY029615Gallusgallus	-----
30	
D50306Rat	-----
NM_053079Musmusculus	-----
XM_007063Homosapiens	-----
AY027496Ovis	-----
U13707Oryctolagusunic	-----
SequencetosubmitGenbak	-----
AY029615Gallusgallus	-----
25	
D50306Rat	-----
NM_053079Musmusculus	-----
XM_007063Homosapiens	-----
AY027496Ovis	-----
U13707Oryctolagusunic	-----
SequencetosubmitGenbak	-----
AY029615Gallusgallus	-----
35	

	D50306Rat	-----AATTGTGTTTCTTGAGACAA	2386
	NM_053079Musmusculus	TGGAGGCATGCACCACCATGCCAGCTATAAATTTTCTTTTAAAGACAG	2620
	XM_007063Homo sapiens	---GGGTTTCACCATGTTGGCCAGG-ATGGTCTCGATCTCTTGACCTGG	2518
	AY027496Ovis	-----	
5	U13707Oryctolagus cuniculus	-----	
	Sequencet osubmit Genbank	-----	
	AY029615Gallus gallus	-----	
10	D50306Rat	GGTATCTCTGTGTAACCCCTGGCTATCCTGGAACTCACTCTATAGACCAGG	2436
	NM_053079Musmusculus	GGATTCTCTGTATAGCCCTGACTGCCCTGGAACTTGCTCTATAGACCAGG	2670
	XM_007063Homo sapiens	TGA---TCTGCCCACCTCGGCCCTGCCAAAGTGTGGGATTACAGGCTTGA	2565
	AY027496Ovis	-----	
	U13707Oryctolagus cuniculus	-----	
	Sequencet osubmit Genbank	-----	
15	AY029615Gallus gallus	-----ATCCTGAGGAAACTCCTGCAGAAATTG	2431
20	D50306Rat	CTGGCCCTCGAACTCACAGATATCTGTCTGCCCTCTGCCCTCCTAAGTACTGG	2486
	NM_053079Musmusculus	CTGGCCCTTGAACTCACAGAGATCTGCCCTGCCCTCTTCTCCTCAAGTACTGG	2720
	XM_007063Homo sapiens	GCTACCGCGCCCGCGCTGAAACGCTATTTTCTAAGCAGCC--AGCAGTGA	2613
	AY027496Ovis	-----GTTTGTGTTGTTTGTAG--AGAAAGTCT	2408
	U13707Oryctolagus cuniculus	-----G-----AAAGTCT	2313
	Sequencet osubmit Genbank	-----	
25	AY029615Gallus gallus	CACCTTTAAAAATGTACCTCAAGCTCAATACCATAGCATTAA-AAAATATTGA	2480
30	D50306Rat	GATTCAAGGCATGTACGGCAACTGCCCCAGCTAAAAATATTATTATAACAT	2536
	NM_053079Musmusculus	GATTTCAGGCATGCACCACAACCTGCCAGCTAAAAATATTATTATAATAT	2770
	XM_007063Homo sapiens	ATCTAAAACCTCTGGAAGAAGTCTTCTGTTTGAAGGCTTATTTAAGCCAC	2663
	AY027496Ovis	TATTTAAAGCGCACAC-ACACGCACACGCACACA-----CAT	2444
	U13707Oryctolagus cuniculus	TATTTAAAACACACAC-ACACACACACACACA-----CAC	2349
	Sequencet osubmit Genbank	-----	
35	AY029615Gallus gallus	AATTGCACCTTGGCACTATTAGACACTCTAAAAAAGATGTATTTT-----TAT	2526

D50306Rat
NM_053079Musmusculus
XM_007063Homo sapiens
AY027496Ovis
U13707Oryctolagus cuniculus
Sequencet osubmit Genbank
AY029615Gallus gallus

GCACCTTCTGGGTTTCTGTTTCTGTTTAAACATACACTTTTCTTTTAAACACTG 2586
GCACCTTCTGG---TTTGTCTTTG-----TTTCTTTTAA-AC TG 2807
ACGTACACACA-----CTGTCTTAGA-----GTACTGTGAGCCACCC 2701
GCACACACACA-----CACTTTTAT-----AAGAGTCCATATC 2478
ACACACTTTTC-----CAACACTG-----ACAGCCTAC---C 2378

ACTGTATTTCAATTTTATAATGTGGAGGGGTGGGAAAAAGGTGTGCCA 2576

5

D50306Rat
NM_053079Musmusculus
XM_007063Homo sapiens
AY027496Ovis
U13707Oryctolagus cuniculus
Sequencet osubmit Genbank
AY029615Gallus gallus

GGCCATTTCTAACATTTCTGCCACAGAAAGTGGAATTAGCTCAGATTAA-- 2634
GGCTGTATCTTACATTTCTGCCACAGAAATGAACCTTAGCTCAGATTAACT 2857
CACATTGGTCATCTTCCCTATCACACAAATGATGTTATTTTGGACTAGCT 2751
TGCCCTGAACCTCTTTCTTAACACACAAATAAAGTTATTTTGGACTAACT 2528
CATGTTAACTCCTTCTCTACCAATGCAAAATGCTGTTATTTTGGACTAACT 2428

AGAAATAGTAATTTGAAGCCAAACCTGTCTGCGTGACCCCTTCTAGCCTCACT 2626

10

D50306Rat
NM_053079Musmusculus
XM_007063Homo sapiens
AY027496Ovis
U13707Oryctolagus cuniculus
Sequencet osubmit Genbank
AY029615Gallus gallus

-----TTTGTGAAAAGGTAAACAGTACTGTGTTTTTTT---TCCTT 2668
T--AATTTTGAAAAGGCAATAGTATTGTTTTTTT-----CT 2890
T--AATTTTGAAAATGGTAACAAAAGTTTCTCTATTCATATCTGTTCACTTCT 2799
TGAATTTTGAATGGTGGCCAAAGCTCCATACGT-----GCATT 2567
T-AATTTTGAACACTGTT-CTATGTTGCTTGTAT-----TC--T 2463

GTTACTTGAAAAGCAGGTCCAC-ATGTGCTTAAATTT-----CTTTTC 2666

15

D50306Rat
NM_053079Musmusculus
XM_007063Homo sapiens
AY027496Ovis
U13707Oryctolagus cuniculus
Sequencet osubmit Genbank
AY029615Gallus gallus

AATGCTCTTA-TGAAAACAATGTTGAA-----TTTACA 2700
AACAGTTTAA-TGAAAACAATATTGAA-----TTTACA 2922
AATACTCTTA-CGAAAACATATCTAAAGGAGGAGGAGGCCAAGGCCAAAA 2848
CGCACACTCTGTGCAACAATGTTAAAGGAGGCAAAAAGTGA---ATGG 2613
AACATCCTTAGGAAAGGCAATGTTAAAGAGGAGGAGGCAATGCCAAAG 2513

TATGTCCTTA---AGAATAATAGGAGAAAG-----GTTTC 2697

30

35

Alignment of Nucleotide Full Length Sequence of Canine and Human

```

Sequence 1: SequencetosubmitGenbank      1840 bp
Sequence 2: XM_007063Homosapiens          3045 bp
Start of Pairwise alignments
Aligning...
Sequences (1:2) Aligned. Score: 85
Guide tree      file created:
[/net/nfs0/voll/production/w3nobody/tmp/305133.88341-239044.dnd]
Start of Multiple Alignment
There are 1 groups
Aligning...
Group 1: Sequences: 2      Score:31290
Alignment Score 10725
15 CLUSTAL-Alignment file created
   [/net/nfs0/voll/production/w3nobody/tmp/305133.88341-239044.aln]

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Your Multiple Sequence Alignment:

20 305133.88341-239044.aln

CLUSTAL W (1.81) multiple sequence alignment

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25 SequencetosubmitGenbank      -----CATCTTCTTC 10
   (SEQ ID NO:7)
   XM_007063Homosapiens      GAATGTCCAAATCACACAGTTTCTTTGGTTATCCCCTGAGCATCTTCTTC 50
   (SEQ ID NO:1)              *****

30 SequencetosubmitGenbank      ATCGTGGTCAATGAGTTCTGTGAAAGATTTTCCCTACTATGGAATGAGAGC 60
   XM_007063Homosapiens      ATCGTGGTCAATGAGTTTTCGAAAGATTTTCCCTACTATGGAATGCGAGC 100
                               ***** ** ***** *****

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SequencetosubmitGenbank XM_007063Homo sapiens	ACTCCTGATTCTGTACTTTCAGACGGTTTCATCGGGTGGGACGATAATCTGT AATCCTGATTCTGTACTTTCACAAATTTTCATCAGCTGGGATGATAACCTGT * * * * *	110 150
SequencetosubmitGenbank XM_007063Homo sapiens	CCACGGCCATCTACACACAGTTTGTGGCTCTGTGCTACCTGACGCCGATC CCACGGCCATCTACACACAGTTTGTGGCTCTGTGCTACCTGACGCCAATT * * * * *	160 200
SequencetosubmitGenbank XM_007063Homo sapiens	CTCGGCGCATGATCGCAGACTCCTGGCTGGGAAAGTTCAAGACAATCTGT CTCGGAGCTCTTATCGCCGACTCGTGGCTGGGAAAGTTCAAGACCAATTGT * * * * *	210 250
SequencetosubmitGenbank XM_007063Homo sapiens	GTCACCTCTCCATTGTCTACACAATTGGACAGGCGGTCACTGCAGTAAGCT GTCGCTCTCCATTGTCTACACAATTGGACAAGCAGTCACCTCAGTAAGCT * * * * *	260 300
SequencetosubmitGenbank XM_007063Homo sapiens	CAATTAATGACCTCACAGACTATAACAAAGATGGAACTCTTGACAACTG CCATTAATGACCTCACAGACCACAAACCATGATGGCACCCCGACAGCCTT * * * * *	310 350
SequencetosubmitGenbank XM_007063Homo sapiens	TCCGTGCATGTGGCAGCTGTCCATGATTGGCCTGGCCCTGATAGCTCTGGG CCTGTGCACGTTGGTGTCTGTCTTGTCTGCGTTTGGTGGAGATCAGTTT * * * * *	360 400
SequencetosubmitGenbank XM_007063Homo sapiens	AACTGGAGGAATAAGCCCTGTGTCTGTGCATTTGGTGGAGACCAGTTT GACTGGAGGAATCAAAACCCCTGTGTCTGCGTTTGGTGGAGATCAGTTT * * * * *	410 450
SequencetosubmitGenbank XM_007063Homo sapiens	AAGAGGGCCAGGAAAAACAAGAAACAGATTCTTTTCCATCTTTTATTG AAGAGGGCCAGGAAAAACAAGAAACAGATTCTTTTCCATCTTTTACTTG * * * * *	460 500
SequencetosubmitGenbank XM_007063Homo sapiens	GCCATTAATGCTGGAAGCTTGATTTCACACTATTGTCACTCCCATGCTCAG GCTATTAATGCTGGAAGTTTGTCTTTCCACAATCATCACCCCATGCTCAG * * * * *	510 550

5	SequencetosubmitGenbank XM_007063Homosapiens	AGTTACGAATGTGGAATTTACAGTCAGAAAAGCTTGTTACCCACTGGCAT AGTTCAACAATGTGGAATTTACAGTAAACAAGCTTGTTACCCACTGGCCT ***** TTGGGGTTCTCTGCTGCTCTCATGGCCCGTATCTCTGATTGTATTTGTCATT TTGGGGTTCTCTGCTGCTCTCATGGCTGTAGCCCTGATTGTGTTTGTCTCTT ***** GGCAGTGGAATGTACAAGAAAGTTTCAGCCCCAGGGTAAATGTCATGGGTAA GGCAGTGGGATGTACAAGAAAGTTCAAGCCACAGGGCAACATCATGGGTAA ***** AGTTGTCAAGTGCATTGGTTTGTCCCTCAAAAAATAGGTTTAGGCACCCGA AGTGCCCAAGTGCATCGGTTTTCGCATCAAAAAATAGATTTAGGCATCGGA *** * GTAAGCAGTTTCCCAAGAGGGAGCAGCTGGCTGGACTGGGCTAAAGAGAAA GTAAGGCATTTCCCAAGAGGGAGCAGCTGGCTGGACTGGGCTAAAGAGAAA **** TACGATGAGCGGCTCATCTCTCAAAATTAAGATGGTCACAAAAGTGATGTT TACGATGAGCGGCTCATCTCCCAAATTAAGATGGTTACGAGGGTGATGTT ***** CTTGATACATCCCACTCCCAATGTTCTGGGCCCTGTTTGACACAGCAGGCT CCTGTATATTCACATCCCAATGTTCTGGGCCCTGTTTGACACAGCAGGCT * **** CCAGGTGGACACTGCAAGCAACAGCTATGAGTGGGAAAATTGGACTTCTT CCAGGTGGACACTGCAGGCAACAACATATGTCCGGGAAAATCGAGCTCTT ***** GAAGTTCAGCCAGATCAGATGCAGACTGTGAATGCCATCTTGATTGTCGT GAAATTCAGCCCCGATCAGATGCAGACCGTGAACGCCATCCTGATCGTAT *** *****
10	SequencetosubmitGenbank XM_007063Homosapiens	*****
15	SequencetosubmitGenbank XM_007063Homosapiens	*****
20	SequencetosubmitGenbank XM_007063Homosapiens	*****
25	SequencetosubmitGenbank XM_007063Homosapiens	*****
30	SequencetosubmitGenbank XM_007063Homosapiens	*****
35	SequencetosubmitGenbank XM_007063Homosapiens	*****

SequencetosubmitGenbank XM_007063Homo sapiens	CATGGTCCCATCATGGATGCCGTGGTGTACCTCTGATTGCAAAATGTG CATGGTCCCGATCTTCGATGCTGTGTACCTCTCATTCGCAAAATGTG ***** *****	1010 1050
SequencetosubmitGenbank XM_007063Homo sapiens	GCTTCAATTTACACCTCCTTGAAGAGGATGACAGTTGGAATGTTCTCTGGCT GCTTCAATTTACACCTCCTTGAAGAGGATGGCAGTTGGCATGGTCTCTGGCC ***** *****	1060 1100
SequencetosubmitGenbank XM_007063Homo sapiens	TCCATGGCCTTCGTGATGGCGGCGATTGTTACAGCTGGAAATGATATAAAC TCCATGGCCTTTGTGGTGGCTGCCATCGTGCAGGTGGAAATCGATAAAC ***** *****	1110 1150
SequencetosubmitGenbank XM_007063Homo sapiens	TCTTCCAGTCTTCCCCAAACAAAATGAAGTCCAAATCAAAGTACTGAATA TCTTCCAGTCTTCCCCAAAGGAAACGAAAGTCCAAATTAAGTTTGAATA ***** *****	1160 1200
SequencetosubmitGenbank XM_007063Homo sapiens	TAGGAAATGGTGCCATGAATGTATCTTTTCTCGAGCGGTGGTGACAGTT TAGGAAACAATACCATGAATATATCTCTTCTCGAGAGATGGTGACACTT ***** *****	1210 1250
SequencetosubmitGenbank XM_007063Homo sapiens	AGCCAAATGAGTCAATCAGATGGATTATGACTTTTGTATGATAGACAAACT GGCCCAATGTCTCAACAAATGCATTTATGACTTTTGTATGATAACAAACT ***** *****	1260 1300
SequencetosubmitGenbank XM_007063Homo sapiens	GACAAATATAAACATTTCTTCCACTGGATCACCAGTCATTCAGTGACTT GACAAAGGATAAACATTTCTTCTCTCGGATCACCAGTCACCTGTAACTG ***** *****	1310 1350
SequencetosubmitGenbank XM_007063Homo sapiens	ATAACTTTGAGCAGGGCCCATCGCCATACCCCTTCTAGTATGGCCCCCAAT ACGACTTCAAGCAGGGCCCAACGCCACACGCTTCTAGTGTGGCCCCCAAT ***** *****	1360 1400
SequencetosubmitGenbank XM_007063Homo sapiens	AATTACCGAGTGGTAAAGGATGGCCTTAACAGAGCCAGAAAAAGGAGA CACTACCGAGTGGTAAAGGATGGTCTTTAACAGAGCCAGAAAAAGGGA ***** *****	1410 1450

SequencetosubmitGenbank XM_007063Homo sapiens	1460 1500	AAATGGAATCAGATTTATAAATAGTCTTAATGAGAGCCTCAACATCACCA AAATGGAATCAGATTTGTAAATACTTTTAAACGAGCTCATCACCATCACAA ***** *
SequencetosubmitGenbank XM_007063Homo sapiens	1510 1550	TGGGCGACAAAAGTTTATGTGAATGTCAACAGTCACAAATGCCAGCGAGTAT TGAGTGGGAAAAGTTTATGCAACATCAGCAGCTACAAATGCCAGCACATAC ** *
SequencetosubmitGenbank XM_007063Homo sapiens	1560 1597	CAGTCTCTTTCTTTGGGCACAAAAAACAATTACAATAAGTTCAACACAACA CAGTCTTTCTCTCTGGCATAAAAGGCTTCACAATAAGCTCAACAG--A ***** *
SequencetosubmitGenbank XM_007063Homo sapiens	1610 1647	GATCTCACAATAATGTACAAAAGTTCTCCAATCATCCAACCTTGAATTG GATTCGGCCACAATGTCAACCTAATTCAATACTTTCTACCTTGAATTG *** *
SequencetosubmitGenbank XM_007063Homo sapiens	1660 1697	GTAGTGCAATACCTATGTAAATCGGAACGCAGAGCAGCTGGCTGCCCTGAA GTAGTGCTTATACCTATATAGTCCAAAGGAAGAAATGACAGCTGCCCTGAA ***** *
SequencetosubmitGenbank XM_007063Homo sapiens	1710 1747	TTGCATATGTTTGAAGATATTTACCCCAACACAGTTAACAATGGCTCTGCA GTGAAGGTGTTTGAAGATATTTACGCCAACACAGTTAACAATGGCTCTGCA ** *
SequencetosubmitGenbank XM_007063Homo sapiens	1760 1797	GATCCCGCAGTACTTCCTCATCACCTGCGGGCGAGGTGTTTCTCTGTCA AATCCCGCAGTATTTTCTCTCACCTGTGGCGAAGTGGTCTTCTCTGTCA ***** *
SequencetosubmitGenbank XM_007063Homo sapiens	1810 1847	CAGGACTGGAGTTCTCATATTTCTCAGGCCCCCTCCAACATGAAGTCGGTG CGGGAATTGGAATTCTCATATTTCTCAGGCTCCTTCCAACATGAAGTCGGTG *
SequencetosubmitGenbank XM_007063Homo sapiens	1840 1897	CTTCAGGCGGATGGCTGCTGACAGTGGCT----- CTTCAGGCGAGGATGGCTGCTGACCGTGGCTGTTGGCAACATCATTTGTGCT ***** *

SequencetosubmitGenbank XM_007063Homo sapiens	----- CATCGTGGCAGGGCAGGCCAGTTCAGCAAAACAGTGGGCCGAGTACATTC 1947
5 SequencetosubmitGenbank XM_007063Homo sapiens	----- TATTTGCCGCGTTGCTTCTGCTCGTCTGTGTAATTTTGTGCCATCATGGCT 1997
10 SequencetosubmitGenbank XM_007063Homo sapiens	----- CGGTTCTATACTTACATCAACCCAGCGGAGATCGAAGCTCAATTTGATGA 2047
15 SequencetosubmitGenbank XM_007063Homo sapiens	----- GGATGAAAAAGAAAAACAGACTGGAAAAAGAGTAACCCATATTTTCATGTCAG 2097
20 SequencetosubmitGenbank XM_007063Homo sapiens	----- GGGCCAATTACAGAAAAACAGATGTGAAGGTCAGGAGGCAAGTGGAGGATG 2147
25 SequencetosubmitGenbank XM_007063Homo sapiens	----- GACTGGGCCCCGAGATGCCCTGACCTCTGTCCCCCAGGTAGCAGGACACTC 2197
SequencetosubmitGenbank XM_007063Homo sapiens	----- CATTGGATGGCCCCCTGATGAGGAAGACTTCAGAAATTGGGAACATAAACCAT 2247
30 SequencetosubmitGenbank XM_007063Homo sapiens	----- GAATGCTATTTTCTTTTCTTTTCTTTTCTTTTCTTTTCTTTTCTTTTCTTTT 2297
35 SequencetosubmitGenbank XM_007063Homo sapiens	----- TTTTTGAGACAGAGTTTGTGCTCTTGTGTCCAGGCTGGAGTGCAATGGCAC 2347

SequencetosubmitGenbank XM_007063Homosapiens	----- GATCTCAGCTCACTGCAACCTCCGCCTCCAGGTTCAAGTAATTCTCTG 2397
5 SequencetosubmitGenbank XM_007063Homosapiens	----- CCTCAGCCTCCCAGAGTGGCTGGGATTAGCGGCATGCACCACGCCCCAG 2447
10 SequencetosubmitGenbank XM_007063Homosapiens	----- CTATTTTGTATTTTAGTAGAGATGGGGTTTCACCATGTTGGCCAGGAT 2497
15 SequencetosubmitGenbank XM_007063Homosapiens	----- GGTCTCGATCTCTTGACCTGGTGATCTGCCCCACCTCGGCCCTGCCAAAGTG 2547
20 SequencetosubmitGenbank XM_007063Homosapiens	----- CTGGGATTACAGGCTTGAGCTACCGCGCCCGCGGTGAACGCTATTTTCT 2597
SequencetosubmitGenbank XM_007063Homosapiens	----- AAGCAGCCAGCAGTGAATCTAAAACTCTGGAAGAAAGTCTTCTGTTGAAA 2647
25 SequencetosubmitGenbank XM_007063Homosapiens	----- GGCTTATTTAAGCCACACGTACACACACTGTCTTAGAGTACTGTGAGCCC 2697
30 SequencetosubmitGenbank XM_007063Homosapiens	----- ACCCACATTTGGTTCATCTTCCCTATCACACAAATGATGTTATTTTGGACT 2747
35 SequencetosubmitGenbank XM_007063Homosapiens	----- AGCTTAATTTTGAAATGGTAACAAAGTTTCCTATTCCTACTGTTCAATT 2797

SequencetosubmitGenbank XM_007063Homo sapiens	----- CTAATACTCTTACGAAAACATATTCTAAAGGAGGCAGGAGCCCAAGGCCAAA 2847
5 SequencetosubmitGenbank XM_007063Homo sapiens	----- AGTGAACGTACAGGTTTGAAAATGGCTGTGATAAGGACCAGCTGGTATTAA 2897
10 SequencetosubmitGenbank XM_007063Homo sapiens	----- CTGATAAACTTTTACCTTTGGGTTTTTTGTATTATTTTCTAGTCCCTAC 2947
15 SequencetosubmitGenbank XM_007063Homo sapiens	----- CTGTGTTTAAATTATGGATAACTCGAAAAGACAGCTCAGGTGAAGGCCAGT 2997
20 SequencetosubmitGenbank XM_007063Homo sapiens	----- AATGATTTTTTTGAAGTTTCAATGGTGTGAAATAAAATTTCTGTTCCTTA 3045

Protein Sequence of Canine

5'3' Frame 2

5 catcttcttcacgtggtcaatgagttctgtgaaagattttcctactatggaatgagagca (SEQ ID NO:8)
I F F I V V N E F C E R F S Y Y G M R A (SEQ ID NO:13)
ctcctgattctgtacttcagacgggttcacatcggtgggacgataatctgtccacggccatc
L L I L Y F R R F I G W D D N L S T A I
10 taccacacgttttggctctgtgctacctgacccgcatcctcggcactgatcgacagac
Y H T F V A L C Y L T P I L G A L I A D
tcctggctgggaaagtccaagacaaatcgtgtcactctccattgtctacacaaattggacag
S W L G K F K T I V S L S I V Y T I G Q
gcgtcactgcagtaagctcaattaatgacctcacagactataacaaaagatggaactcct
A V T A V S S I N D L T D Y N K D G T P
15 gacaatctgtccgtgcattgtggcactgtccatgattggccctggccctgatagctctggga
D N L S V H V A L S M I G L A L I A L G
actggaggaataaagccctgtgtgtctgcatcttggtggagaccagtttgaagagggccag
T G G I K P C V S A F G G D Q F E E G Q
gaaaaacaaagaacacagattctttccatcttttatttggccattaatgctggaagcttg
E K Q R N R F F S I F Y L A I N A G S L
20 atttccactattgtcactcccatgctcagagttcacgaatgtggaatttacagtcagaaa
I S T I V T P M L R V H E C G I Y S Q K
gcttgttaccactggcatttggggttcctgctgctctcatggtccgtatctctgattgta
A C Y P L A F G V P A A L M A V S L I V
25 ttgtcattggcagtggaatgtacaaagaagtttcagccccagggtaatgtcatgggtaaa
F V I G S G M Y K K F Q P Q G N V M G K
gttgcgaagtgcattggttttggccctcaaaaatagggttaggcaccggagtaagcagttt
V V K C I G F A L K N R F R H R S K Q F
cccaagaggagcactggctggactgggttaaagagaaatacagatgagcggctcatctct
P K R E H W L D W A K E K Y D E R L I S
30 caaattaagatggtcacaaaagtgatgttcttgtacatccccactccccaatgttctgggccc
Q I K M V T K V M F L Y I P L P M F W A

ctgtttgaccagcagggtccagggtggacactgcaagcaacagctatgagtgggaaaaatt
 L F D Q Q G S R W T L Q A T A M S G K I
 ggacttcttgaagttcagccagatcagatgcagactgtgaatgccatcttgattgtcgtc
 G L L E V Q P D Q M Q T V N A I L I V V
 5 atggtcccatcatggatgccgtggtgtaccctctgattgcaaaatgtggcttcaatttc
 M V P I M D A V V Y P L I A K C G F N F
 acctccttgaagaggtgacagttggaatgttcctggcttccatggccttcgtgatggcg
 T S L K R M T V G M F L A S M A F V M A
 gcgattgttcagctggaaaattgataaaaactcttccagctcttccccaaaaaatgaagtc
 10 A I V Q L E I D K T L P V F P K Q N E V
 caaatcaaaagtactgaatataggaaaatggtgccatgaatgtatcttttccctggagcgggtg
 Q I K V L N I G N G A M N V S F P G A V
 gtgacagtttagccaaaatgagtcfaatcagatggatttatgactttttgatgtagacaaaactg
 V T V S Q M S Q S D G F M T F D V D K L
 15 acaagtataaaacatttctccactggatcacccagtcattccagtgacttataactttgag
 T S I N I S S T G S P V I P V T Y N F E
 cagggccatcgccatacccttcttagtatggcccccaataattaccgagtggtaaaggat
 Q G H R H T L L V W A P N N Y R V V K D
 ggccttaaccagaagccagaaaaaggagaaaaatggaatcagatttataaatagtccttaat
 20 G L N Q K P E K G E N G I R F I N S L N
 gagagcctcaacatcaccatggcgacaaaagtttatgtgaatgtcacccagtcacaaatgcc
 E S L N I T M G D K V Y V N V T S H N A
 agcgagtatcagttcttcttcttgggcacaaaaaacattacaataagttcaacacaacag
 S E Y Q F F S L G T K N I T I S S T Q Q
 25 atctcacaaaattgtacaaaagttctccaatcatccaaccttgaatttggtagtgcatat
 I S Q N C T K V L Q S S N L E F G S A Y
 acctatgtaatcggaacgcagagcactggctgccctgaattgcatatgtttgaagatat
 T Y V I G T Q S T G C P E L H M F E D I
 tcacccaacacagttaacatggctctgcagatcccgcagtagtctcctcatcacctgcggc
 30 S P N T V N M A L Q I P Q Y F L I T C G
 gaggtggtttctctgtcacaggactggagttctcatattctcaggccccctccaacatg
 E V V F S V T G L E F S Y S Q A P S N M

aagtcggtgcttcaggcgggatggctgctgacagtggtggtggcaacatcattgtgct
 K S V L Q A G W L L T V A C W Q H H C A
 cattgtggcaggagcagccagttcagtgacagtggtggtgaatacatcctatttcgggc
 H C G R S R P V Q - T V G - I H P I C G
 attgcttctggtgctgtgtaataattgcatcatcagggcccggttttacacttacgtcaa
 I A S G C L C N I C H H G P V L H L R Q
 tccagcagagattg
 S S R D

5

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5'3' Frame 2 (SEQ ID NO:13)

IPFIVVNEFCERFSYYGMRALLILYFRRFIGWDDNLSTAIYHTFVALCYLTPILGALIA
 SWLGKFKTI VSLSI VYTI GAVTAVSSINDLTDYKDGTPDNL SVHVALSMIGLALIALG
 TGGIKPCVSAFGGDQFEEGQEKQRNRFSSIFYLA INAGSLISTIVTPMLRVHECGIYSQK
 ACYPLAFGVPAALMAVSLIVFVIGSGMYKKFPQGNVVMGKVVKCIGFALKNRFRHRSKQF
 PKREHWLDWAKEKYDERLISQIKMVTKMFLYIPLPMFWALFDQQGSRWTLQATAMSGKI
 GLLLEVQPDQMOTVNAIILVVMVPIMDAVVYPLIAKCGFNFTSLKRMTVGMFLASMAFVMA
 AIVQLEIDKTLPVFPKQNEVQIKVLNIGNGAMNVSPGAVTVSQMSQSDGFMTFDVKL
 TSINISSTGSPVIPVTYNFEQGHRTLLVWAPNNYRVVKDGLNQKPEKGENGIRFINSLN
 ESLNITMGDKVYVNVTSHNASEYQFFSLGTKNITISSTQQISQNC TKVLQSSNLEFGSAY
 TYVIGTQSTGCP ELHMFEDI SPNTVNMA LQIPQYFLITCGEVVFSVTGLEFSYSQAPS NM
 KSVLQAGWLLTVACWQHHC AHCGRSRPVQ-TVG-IHPICG IASGCLCNI CHHGPVLHLRQ
 SSRD

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Multiple Alignment of Amino-Acid Sequences

Sequence 1: Caninesubmitted 662 aa
 Sequence 2: XM_007063HomosapiensProteinSeq 706 aa
 Sequence 3: D50306RatProteinSequence 710 aa
 Sequence 4: NM_053079MusmusculusProteinSeq 709 aa
 Sequence 5: AY027496Ovis 707 aa
 Sequence 6: U13707OryctolagusCunicProteins 707 aa
 Sequence 7: AY029615GallusgallusProteinSeq 714 aa

10 Start of Pairwise alignments

Aligning...

Sequences (1:2) Aligned. Score: 76
 Sequences (2:3) Aligned. Score: 84
 Sequences (3:4) Aligned. Score: 91
 Sequences (4:5) Aligned. Score: 80
 Sequences (1:3) Aligned. Score: 77
 Sequences (2:4) Aligned. Score: 83
 Sequences (3:5) Aligned. Score: 82
 Sequences (4:6) Aligned. Score: 76
 Sequences (1:4) Aligned. Score: 75
 Sequences (2:5) Aligned. Score: 82
 Sequences (3:6) Aligned. Score: 77
 Sequences (4:7) Aligned. Score: 63
 Sequences (1:5) Aligned. Score: 77
 Sequences (2:6) Aligned. Score: 80
 Sequences (1:6) Aligned. Score: 72
 Sequences (3:7) Aligned. Score: 64
 Sequences (5:6) Aligned. Score: 77
 Sequences (1:7) Aligned. Score: 60

25 Sequences (6:7) Aligned. Score: 61Sequences (2:7) Aligned. Score: 63
 Sequences (5:7) Aligned. Score: 64

Guide tree file created:

[/net/nfs0/voll/production/w3nobody/tmp/936042.678539-441485.dnd]

Start of Multiple Alignment

There are 6 groups

Aligning...

5	Group 1:	Sequences:	2	Score:14016
	Group 2:	Sequences:	2	Score:14858
	Group 3:	Sequences:	4	Score:13893
	Group 4:	Sequences:	5	Score:14022
10	Group 5:	Sequences:	6	Score:12718
	Group 6:	Sequences:	7	Score:12338

Alignment Score 68091

CLUSTAL-Alignment file created

[/net/nfs0/voll/production/w3nobody/tmp/936042.678539-441485.aln]

15 Your Multiple Sequence Alignment:

936042.678539-441485.aln

CLUSTAL W (1.81) multiple sequence alignment

20

	XM_007063HomosapiensProteinSeq (SEQ ID NO:14)	---MSKSHS-----FFGYPLSIFFIVVNEFCERFSYYGMRALLILLYFTNF	44
	U13707OryctolagusCunicProteins (SEQ ID NO:18)	-MGMSKSLS-----CFGYPLSIFFIVVNEFCERFSYYGMRALLILYFRNF	44
25	D50306RatProteinSequence (SEQ ID NO:15)	-MGMSKSRG-----CFGYPLSIFFIVVNEFCERFSYYGMRALLVL YFRNF	44
	NM_053079MusmusculusProteinSeq (SEQ ID NO:16)	-MGMSKSRG-----CFGYPLSIFFIVVNEFCERFSYYGMRALLVL YFRNF	44
	AY027496Ovis (SEQ ID NO:17)	-MGMSVPKS-----CFGYPLSIFFIVVNEFCERFSYYGMRALLILYFQRF	44
30	Caninesubmitted (SEQ ID NO:13)	-----IFFIVVNEFCERFSYYGMRALLILYFRRF	29
	AY029615GallusgallusProteinSeq (SEQ ID NO:19)	MAAKSKSGRSVPNCFGYPLSIFFIVINEFCERFSYYGMAVLVL YFKYF	50
		*****;*****;*:*** *	*

	XM_007063Homo sapiensProteinSeq	I SWDDNLSTAIYHTFVALCYLTPILGALIASWLGKFTIVSLSI VYTIG	92
	U13707Oryctolagus cuniculusProteins	IGWDDNLS TVIYHTFVALCYLTPILGALIA DAWLGKFTIVWL SI VYTIG	94
	D50306RatProteinSequence	LGWDDDLSTAIYHTFVALCYLTPILGALIASWLGKFTIVSL SI VYTIG	94
5	NM_053079Mus musculusProteinSeq	LGWDDNLS TAIYHTFVALCYLTPILGALIASWLGKFTIVSL SI VYTIG	94
	AY027496Ovis	L GWNDNLGT AIYHTFVALCYLTPILGALIASWLGKFTIVSL SI VYTIG	94
	Caninesubmitted	IGWDDNLS TAIYHTFVALCYLTPILGALIASWLGKFTIVSL SI VYTIG	79
10	AY029615GallusgallusProteinSeq	LRWDDNFSTAIYHTFVALCYLTPILGALIASWLGKFTIVSL SI VYTIG	100
		: *: : : *. *****:*****:***** ***** *	
	XM_007063Homo sapiensProteinSeq	QAVTSVSSINDLTDNHDGTPD SLPVHVVL SL IGLALIALGTGGIKPCVS	142
	U13707Oryctolagus cuniculusProteins	QAVTSLSSVNELTDNNHDGTPD SLPVHVAVCMIGLLLI ALGTGGIKPCVS	144
	D50306RatProteinSequence	QAVISVSSINDLTDDHDGSPNNPLPHVALSMIGLALIALGTGGIKPCVS	144
	NM_053079Mus musculusProteinSeq	QAVISVSSINDLTDDHNGSPD SLPVHVALS MVGLALIALGTGGIKPCVS	144
15	AY027496Ovis	QVVI AVSSINDLTDFNHGDGPNNISVHV ALSMIGLVLI ALGTGGIKPCVS	144
	Caninesubmitted	QAVTA VSSINDLT DYNKDGTPD NLSVHV ALSMIGLALIALGTGGIKPCVS	129
	AY029615GallusgallusProteinSeq	QAVMAVSSINDMTDQNDRGNPDNI AVHIALSMTGLILIALGTGGIKPCVS	150
		..:*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*	
20	XM_007063Homo sapiensProteinSeq	AFGGDQFE EGQE KQRNRFFSIFYLAINAGS LLSTIITPMLRVQQGIHSK	192
	U13707Oryctolagus cuniculusProteins	AFGGDQFE EGQE KQRNRFFSIFYLAINAGS LLSTIITPMVRVQQGIHV K	194
	D50306RatProteinSequence	AFGGDQFE EGQE KQRNRFFSIFYLAINAGS LLSTIITPILRVQQGIHS Q	194
	NM_053079Mus musculusProteinSeq	AFGGDQFE EGQE KQRNRFFSIFYLAINAGS LLSTIITPILRVQQGIHS Q	194
	AY027496Ovis	AFGGDQFE EGQE KQRNRFFSIFYLAINAGS LLSTIITPMLRVQVC GIHSK	194
25	Caninesubmitted	AFGGDQFE EGQE KQRNRFFSIFYLAINAGS LI STIVT PMLRVHEC GIYSQ	179
	AY029615GallusgallusProteinSeq	AFGGDQFE EH QE KQRS RFFSIF YLSINAGS LI STIITPILRAQEC GIHSR	200
		***** ***. *****:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*	
	XM_007063Homo sapiensProteinSeq	QACYPLAFGVPAALMA VALIVFVLGSGMYKKFKPQGNIMGKVAKICGF AI	242
	U13707Oryctolagus cuniculusProteins	QACYPLA FGI PAI LMAVSLIVFI IGSGMYKKFKPQGNILSKVVKCICFAI	244
30	D50306RatProteinSequence	QACYPLA FGVPAALMA VALIVFVLGSGMYKKFPQPGNIMGKVAKICGF AI	244
	NM_053079Mus musculusProteinSeq	QACYPLA FGVPAALMA VALIVFVLGSGMYKKFPQPGNIMGKVAKICGF AI	244
	AY027496Ovis	QACYPLA FGVPAALMA VSLIVFVIGSGMYKKVPQPGNIMSKVARCIGF AI	244
	Caninesubmitted	KACYP LA FGVPAALMA VSLIVFVIGSGMYKKFPQPGNVMMKVVKCIGF AL	229
35	AY029615GallusgallusProteinSeq	QQCYPLA FGVPAALMA VSLIVFV IAGSGMYKKVPQPGNIMVRVCKICGF AI	250
		: *****:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*:~*	

[illegible]

XM_007063Homo sapiens ProteinSeq	AGQFSKQWAEYILFAALLLVVCVIFAIMARFYTYINPAEIEAQFDEDEKK	686
U13707Oryctolagus cunic ProteinS	AGQINKQWAEYILFAALLLVVCVIFAIMARFYTYVNPAAIEAQFEEDEKK	687
D50306Rat ProteinSequence	AGHFDKQWAEYVLFASLLLVVCIIIFAIMARFYTYINPAEIEAQFDEDEKK	690
NM_053079Mus musculus ProteinSeq	AGHFPKQWAEYILFASLLLVVCVIFAIMARFYTYINPAEIEAQFDEDEKK	689
AY027496Ovis	AGQFSEQWAEYVLFASLLLVVCIIIFAIMARFYTYVNPAAIEAQFDEDDKE	687
Caninesubmitted	SRPVQ-TVG-----IHPICGIASGCLCNI CHGPPVLHLRQSSRD	662
AY029615Gallus gallus ProteinSeq	ASKLSEQWAEYVLFALLFAVCIIFAVMAYFYTYTDPNEVEAQLDEEEKK	694

10

XM_007063Homo sapiens ProteinSeq	NRLEKSNPYFMMSGANSQKQM	706
U13707Oryctolagus cunic ProteinS	KNPEKNDLYPSVAPVSQTQM	707
D50306Rat ProteinSequence	KGVGKENPYSSLEPVSQTNM	710
NM_053079Mus musculus ProteinSeq	KGIGKENPYSSLEPVSQTNM	709
AY027496Ovis	DDLEKSNPYAKLDFVSQTQM	707
Caninesubmitted	-----	
AY029615Gallus gallus ProteinSeq	KQIKQDPDLHGKSEAVSQM	714

15

Alignment of Amino-Acid Sequences for Canine and Human

Sequence format is Pearson

Sequence 1: XM_007063HomoapiensProteinSeq 706 aa
 Sequence 2: Caninesubmittedclone37 662 aa

Start of Pairwise alignments

Aligning...

Sequences (1:2) Aligned. Score: 76

Guide tree file created:

10 [/net/nfs0/voll/production/w3nobody/tmp/789481.229198-238519.dnd]

Start of Multiple Alignment

There are 1 groups

Aligning...

Group 1: Sequences: 2 Score:12826

15 Alignment Score 3129

CLUSTAL-Alignment file created

[/net/nfs0/voll/production/w3nobody/tmp/789481.229198-238519.aln]

Your Multiple Sequence Alignment:

789481.229198-238519.aln

CLUSTAL W (1.81) multiple sequence alignment

25

XM_007063HomoapiensProteinSeq MSKSHSFFGYPLSIFFIIVNEFCERFSYGMRAILILYFTNFISWDDNLS 50
 (SEQ ID NO:14)
 Caninesubmittedclone37
 (SEQ ID NO:13)

-----IFFIIVNEFCERFSYGMRAILILYFRRFIGWDDNLS 37
 *****;*****

30

XM_007063HomoapiensProteinSeq
 Caninesubmittedclone37

TAIYHTFVALCYLTPILGALIADSWLGKFKTIVSLSIYVTIGQAVTSVSS 100
 TAIYHTFVALCYLTPILGALIADSWLGKFKTIVSLSIYVTIGQAVTSVSS 87
 *****;*****

XM_007063HomosapiensProteinSeq
Caninesubmittedclone37
SAYTYIVQRKNDSCPEVKVFEDISANTVNMALQIPQYFLITCGEVVFSVT 599
SAYTYVIGTQSTGCPELHMFEDISPNTVNMALQIPQYFLITCGEVVFSVT 587
*****: :.***:::*****.*****;*****

5

XM_007063HomosapiensProteinSeq
Caninesubmittedclone37
GLEFSYSQAPSNMKSVLQAGWLLTVAVGNIIVLIVAGAGQFSKQWAEYIL 649
GLEFSYSQAPSNMKSVLQAGWLLTVACWQHHAHCAHGRSRPVQ-TVG---- 632
*****: :.***:::*****.*****;*****

10

XM_007063HomosapiensProteinSeq
Caninesubmittedclone37
FAALLLVVCVIFAIMARFYTYINPAEIEAQFDEDEKKNRLEKSNPYFMSG 699
-----IHPICGIASGCLCNICHGHPVLHLRQSSRD----- 662
: *. * : * : : : : .

15

XM_007063HomosapiensProteinSeq
Caninesubmittedclone37
ANSQKQM 706

After analyzing the protein sequence and performing alignment with other species, the underlined, italicized was removed for submission to Genbank.

5 **Sequence to submit to Genbank (SEQ ID NO:7)**

catcttctcatcgtggtcaatgagttctgtgaaagatttctactatggaatgagagcactcctgattctgtacttcagacgg
ttcatcgggtgggacgataatctgtccacggccatctaccacacgtttgtggctctgtgctacctgacgccgatcctcggc
gcactgatcgagactcctggctgggaaagtcaagacaatcgtgcactctccattgtctacacaattggacaggcggtc
actgcagtaagctcaattaatgacctcacagactatacaaagatggaactcctgacaatctgtccgtgcagtgtggcactgt
10 ccatgattggcctggccctgatagctctgggaactggaggaataaagccctgtgtgtctgcatttggaggagaccagtttg
aagagggccaggaaaaacaagaacagattctttccatctttatttggccattaatgctggaagcttgattccactattg
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ctcatggccgtatctctgattgtatttgcattggcagtggaaatgtacaagaagtttcagccccagggtaatgcatgggttaa
agttgtcaagtgcattggtttgcctcaaaaataggttaggcaccggagtaagcagtttccaagaggagcactggct
15 ggactgggctaagagaaatagcatgagcggctcatctctcaaattaagatggtcacaaaagtgtgttctgtacatccc
actcccaatgttctgggccctgtttgaccagcagggtccaggtggacactgcaagcaacagctatgagtgggaaaattg
gacttctgaagttcagccagatcagatgcagactgtgaatgccatcttgattgtcgtcatgggtcccatcatggatgccgt
gggtgtaccctctgattgcaaatgtggcttcaattcacctccttgaagaggatgacagttggaatgttctggcttccatgg
ccttcgtgatggcggcgattgttcagctggaaattgataaaactcttcagcttccccaaacaaaatgaagtccaaatcaa
20 agtactgaatataggaaatggtgccatgaatgtatctttctggagcgggtggtagacagttagccaaatgagtcaatcagat
ggatttatgacttttgatgtagacaaactgacaagtataaacatttctccactggatcaccagtcattccagtacttataact
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ccagaaaaaggagaaaatggaatcagattataaatagtcttaatgagagcctcaacatcaccatgggcgacaaagtatat
gtgaatgtcaccagtcacaatgccagcgagtatcagttcttttcttgggcacaaaaacattacaataagttcaacacaac
25 agatctcacaaaattgtacaaaagtttccaatcatccaaccttgaatttgtagtgcataacatgtaatcggaacgcag
agcactggctgccctgaattgcataatgttgaagatattcacccaacacagttaacatggctctgcagatcccgagctactt
cctcatcacctgcggcgaggtggtttctctgtcacaggactggagttctcatattctcagggccccctcaacatgaagtcg
gtgcttcaggcgggatggctgctgacagtggt

30 **Canine PepT1 Nucleotide Sequence (SEQ ID NO:20)**

atgggcgatgtccaagtcatatggttgctttggttacccttgagcatcttcttcacgtggtcaatgagttctgtgaaagattt
cctactatggaatgagagcactcctgattctgtacttcagacgggtcatcgggtgggacgataatctgtccacggccatcta

ccacacgtttgtggctctgtgtacctgacgccgatcctcggcgcactgatcgagactcctggctgggaaagtcaaga
 caatcgtgtcactctccattgtctacacaattggacaggcggcactgcagtaagctcaattaatgacctcagactataa
 caaagatggaactcctgacaatctgtccgtgcatgtggcactgtccatgattggcctggccctgatagctctgggaactgg
 aggaataaagccctgtgtgtctgcatttgggtgagaccagtttgaagagggccaggaaaaacaaagaaacagattcttt
 5 ccatcttttatttggccattaatgctggaagcttgattccactattgtcactcccatgctcagagttcacgaatgtggaatttac
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 25

Canine PepT1 Amino Acid Sequence (SEQ ID NO:21)

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 35 GAVVTVSQMSQSDGFMTFDVDKLT SINISSTGSPVIPVTYNFEQGHRHTLLV
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 AGWLLTVAVGNIIVLIVAGAGQFSEQWAEYILFAALLLVVCVIFAIMARFYT
 40 YVNP AEIEAQFDDDEKKNLEKMNVYSTVTPVSQTQM

All publications, patents and patent documents are incorporated by reference herein, as though individually incorporated by reference. The invention has been

described with reference to various specific and preferred embodiments and techniques. However, it should be understood that many variations and modifications may be made while remaining within the scope of the invention.

2020-03-04 10:40:00